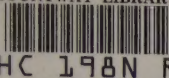


COUNTWAY LIBRARY



HC 198N R

Francis Cooley Hall

Harvard Medical School '17

INFECTION AND RESISTANCE



THE MACMILLAN COMPANY
NEW YORK • BOSTON • CHICAGO
DALLAS • ATLANTA • SAN FRANCISCO

MACMILLAN & CO., LIMITED
LONDON • BOMBAY • CALCUTTA
MELBOURNE

THE MACMILLAN CO. OF CANADA, LTD.
TORONTO

INFECTION AND RESISTANCE

AN EXPOSITION OF THE BIOLOGICAL PHENOMENA
UNDERLYING THE OCCURRENCE OF INFECTION
AND THE RECOVERY OF THE ANIMAL BODY
FROM INFECTIOUS DISEASE

BY

HANS ZINSSER, M.D.

Professor of Bacteriology at the College of Physicians and Surgeons, Columbia University,
New York. Formerly Professor of Bacteriology and Immunity
at Stanford University, California

WITH A CHAPTER ON

COLLOIDS AND COLLOIDAL REACTIONS

BY

PROFESSOR STEWART W. YOUNG

Department of Chemistry, Stanford University

New York

THE MACMILLAN COMPANY

1914

BOSTON MEDICAL LIBRARY
IN THE
FRANCIS A. COUNTWAY
LIBRARY OF MEDICINE

COPYRIGHT 1914
BY THE MACMILLAN COMPANY

Set up and electrotyped. Published October, 1914

TO

A. Z.

THIS BOOK IS AFFECTIONATELY

DEDICATED BY HIS

SON

PREFACE

INFECTIOUS disease, biologically considered, is the reaction which takes place between invading micro-organisms and their products, on the one hand, and the cells and fluids of the animal's body on the other. The disease is the product of two variable factors, each of them to a certain extent amenable to analysis, and it is self-evident that no true understanding of this branch of medicine is possible without a knowledge of the biological principles which laboratory study has revealed.

For the purpose of helping to render such knowledge easily accessible this book was written. While it is hoped that it may prove useful to the practitioner and laboratory worker, it is intended primarily for the undergraduate medical student. To many it will seem that the subject in general and our method of treatment especially are too technical and difficult for this purpose. Our own experience contradicts this. During the past three years the writer has had the opportunity to deliver lectures and to give laboratory courses on this subject to medical students of 2d, 3d, and 4th-year classes at the Stanford and Columbia Universities. It has been a pleasant experience to find the medical student eager for the opportunity to obtain this knowledge and, under the present increased requirements for preliminary training at our best schools, fully capable of assimilating it. It is not a good plan to attempt too extensively to simplify material that, in its close analysis, presents complex phenomena and intricate reasoning. For this reason no attempt has been made to write an A B C of immunity as a quick road to comprehension. No true insight into any branch of medicine or, for that matter, into any other science, can be attained without a certain amount of labor; however the concepts of this subject are, indeed, relatively simple after the first principles have been mastered, and the writer has attempted, therefore, at the risk of seeming pedantic in places, to treat the subject critically, separating strictly those data which may be accepted as fact from those in which legitimate differences of opinion prevail.

As far as was feasible every chapter has been written as a separate unit. This has necessitated occasional repetition, but, it is hoped, will add considerably to clearness of presentation in each individual subject. Theories have been discussed with as little prejudice as the possession of a personal opinion in many cases has permitted.

The chapter on Colloids was written especially for the book by Prof. Stewart W. Young, of Stanford University. Since so many analogies between serum reactions and those taking place between colloidal substances generally have been observed, it has seemed best to devote this chapter entirely to the elucidation of the principles governing colloidal reactions, so that its contents may be utilized as explanatory of the many allusions made to colloids in the rest of the text.

All available sources of information have been freely used. In the large majority of cases we have had access to the original papers and monographs. However, we acknowledge much aid from careful reading of the admirable summaries, written by acknowledged authorities, in the works edited by Kolle and Wassermann, and by Kraus and Levaditi. Similar acknowledgment is made to equally important sources in Weichhardt's *Jahresbericht*, the *Bulletins* of the Pasteur Institute, and in such text-books as those of Paul Theo. Müller, Emery, Adami, Gideon Wells, Marx, Dieudonné, and others. It is needless to acknowledge the use of such classics as that of Metchnikoff or of the many critical writings of Bordet and of Ehrlich—masters who have helped to shape the thoughts of all men working in this field.

The writer takes pleasure in acknowledging many helpful suggestions from his associates, Drs. Hopkins and Ottenberg, and much aid, in the verification of references, from Mr. Walter Bliss, Fellow in the Department of Bacteriology.

CONTENTS

| | PAGE |
|--|------|
| CHAPTER I.—INFECTION AND THE PROBLEM OF VIRULENCE | 1 |
| <p>Scope of subject. Conception of infection. Attributes of pathogenic microorganisms. Forms of infection. Influences of biological adaptation. Classification of parasites on the basis of invasive properties. Factors which determine the power to invade. Fluctuations in virulence. How microorganisms defend themselves against destruction. Serum fastness, arsenic fastness, capsule formation, inagglutinability, etc. Resistance to phagocytosis. Development of offensive properties on the part of bacteria. Specificity of different infections. Chronic septicæmia. "Sub-infection." Selective lodgment in tissues. Localization and generalization. Incubation time.</p> | |
| CHAPTER II.—BACTERIAL POISONS | 23 |
| <p>Part played by bacterial poisons in clinical manifestations. Pto- maines. Importance of ptomaines in disease. True toxins or exo- toxins. Endotoxins. Chemotactic bacterial extracts. Basic proper- ties of true toxins. Other substances biologically similar to them. Analogy to enzymes. Snake venoms. Incubation time of toxins. Conception of antitoxins. Work of Vaughan. Researches of Fried- berger. Absorption of toxins. Selective action of toxins. Distribu- tion of tetanus poison. Causes underlying selective action in general. Injury done during the excretion of toxins. Union of toxins with sus- ceptible cells. Importance of cell lipoids.</p> | |
| CHAPTER III.—OUR KNOWLEDGE CONCERNING NATURAL IMMUNITY, AC- QUIRED IMMUNITY AND ARTIFICIAL IMMUNITY | 49 |
| <p>The struggle between the infectious agent and the defensive forces of the body. External defenses. Skin secretions. Natural vs. arti- ficially acquired immunity. Species immunity. Racial immunity. Dif- ference between individuals. Inheritance of natural immunity. Im- munity resulting from an attack of the disease. Jenner and smallpox. Pasteur's work with chicken cholera. Active immunization. Passive immunization. Pasteur's studies on anthrax. Different methods of con- ferring active immunity. Methods of obtaining bacterial extracts. De- velopment of our knowledge of passive immunization. Early attempts. Behring and his collaborators. Ehrlich's work on ricin. Snake venom. Specificity.</p> | |
| CHAPTER IV.—THE MECHANISM OF NATURAL IMMUNITY AND THE PHE- NOMENA FOLLOWING UPON ACTIVE IMMUNIZATION | 78 |
| <p>Investigations on problems of inflammation. Metchnikoff's earlier studies. Concentration of attention upon the properties of the blood. Grohman's work. Early opposition of cellular and humoral points of view. Buchner. Nuttall. Earlier arguments brought forward by the two schools. Behring's summary of the situation at this time. Phenomena following upon active immunization. Earlier theories. Exhaustion theory. Retention theory. Alkalinity theory. Osmotic theory. Discovery of specific antibodies by Behring and collaborators. Ehrlich's study on ricin. Antitoxins. Pfeiffer's discovery of lysins. Agglutinins. Precipitins. Opsonins. Tropins. Conception of anti- bodies as a whole. Generalization of the facts discovered in the case</p> | |

of bacteria. Hæmolysins. Cytotoxins. Hæmagglutinins. Precipitins to unformed proteins. Conception of an antigen. Nature of antigens. Analogy of immunity with drug tolerance.
Origin of antibodies.

CHAPTER V.—TOXIN AND ANTITOXIN 104

Nature of the reaction. Earlier views. Calmette's work on snake poison. Filtration experiments. Morgenroth's work on HCl-toxin modifications. Ehrlich's ricin neutralization. Development of the neutralization ideas by Ehrlich and Behring. Conception of antitoxin unit. Instability of toxin. Ehrlich's experiments. The conceptions of M.L.D., L_0 and L_+ doses. Discrepancy between L_0 and L_+ . Toxoids and toxons. Method of partial absorption. Toxin spectrum. Opinions of Arrhenius & Madsen. Bordet's opinion. The Danyz effect. THE SIDE CHAIN THEORY. Early work of Knorr. Ehrlich's analogy of cell with chemical substances. Analogy with ferments. Weigert's law of overcompensation. Antibodies and cell receptors. Theoretical conclusions drawn from work with tetanus poison. Importance of lipoidal substances in brain tissue.

CHAPTER VI.—BACTERICIDAL PROPERTIES OF BLOOD SERUM, CYTOLYSIS AND SENSITIZATION 134

The phenomenon of bacteriolysis and the bactericidal effect. Hæmolysis. The mechanism of cytolysis. Amboceptor or sensitizer? The complement or alexin. Iso-antibodies. Discussion of views of Ehrlich and Bordet. Multiplicity of antibodies. Multiplicity of complement or alexin. Anti-antibodies. Neisser and Wechsberg phenomenon of complement deviation. Quantitative relation between complement and amboceptor. Conglutinins.

CHAPTER VII.—DEVELOPMENT OF OUR KNOWLEDGE CONCERNING COMPLEMENT OR ALEXIN. COMPLEMENT FIXATION 168

Origin of alexin. Microcytase and Macrocytase. Anti-lysins. Alexin in œdema fluids, etc. The question of alexin in the circulating blood. Alexin and the thyroid. Alexin and the liver. Chemical nature of complement and alexin. Cobra-lecithid. Enzyme-like nature of alexin. Filtration of alexin. Complement or alexin splitting. Return to activity of inactivated alexin on standing. Inactivation by shaking. ALEXIN FIXATION. Bordet-Gengou experiments. Theoretical explanation of these facts. Albuminolysins of Gengou. Alexin fixation by precipitates. Views of earlier writers. Nicoll's view. Writer's opinion. Conception of "Bordet-antibody." Nonspecific alexin fixation. Importance of lipoids. Fixation by unsensitized cells, substances in suspension. Anticomplementary properties of serum.

CHAPTER VIII.—PRACTICAL APPLICATIONS OF COMPLEMENT-FIXATION METHOD. THE WASSERMANN REACTION 198

Historical. Early work on monkeys. First use on human beings. Theories of the Wassermann reaction. Methods of preparing antigen. Titration of antigen. Titration of hæmolytic sensitizer. Alexin titration. Performance of the test. Noguchi's modification. Modifications of Bauer, Stern and others. Results of test. Reliability. Spinal fluid, etc. COMPLEMENT OR ALEXIN FIXATION FOR THE DETERMINATION OF UNKNOWN PROTEIN. Neisser-Sachs method. Principles of the method. Performance of test. COMPLEMENT-FIXATION TESTS IN DIAGNOSIS OF MALIGNANT NEOPLASMS. Historical. Von Dungern's method. Results obtained. Complement-fixation in glanders. Complement-fixation in gonococcus infections.

CHAPTER IX.—THE PHENOMENON OF AGGLUTINATION 218

Discovery. Applications of clinical methods. Clinical usefulness. Relation to motility. Passive rôle of bacteria. Bordet's discovery of

CONTENTS

xi

PAGE

the importance of electrolytes. Nature of agglutininogen. Alterations by heat. Alterations in agglutinability. Reasons for agglutinability. Specificity. Biological relations between bacteria parallel to agglutinins. Castellani's method of absorption. Normal agglutinins. Agglutinoids. Inhibition zones. Bordet's views. "Two-phase" theory. Physical interpretation. The work of Neisser and Friedemann. Acid agglutination. Iso-agglutinins.

CHAPTER X.—THE PHENOMENA OF PRECIPITATION 248

Discovery. Bacterial filtrates. Expansion of principle to proteins in general. Nature of precipitinogen. Specificity. Quantitative relations in the reaction. Practical uses. Nuttall's studies on precipitins and historical relationship. Forensic uses of the test. Performance of the test as advised by Uhlenhuth. Influence of heat upon precipitinogen. Organ specificity. Ehrlich's view of the nature of the reaction. Physical views of the reaction. Presence of precipitinogen and precipitin in same serum. Analogy with colloids of known constitution.

CHAPTER XI.—PHAGOCYTOSIS. CHEMOTAXIS. 272

Early investigations. Metchnikoff's first studies. Phagocytosis in lower animals. Its significance. Importance in the development from the larva to the adult. Its importance in resorption of degenerated cells. Varieties of phagocytosis. Giant cells. Leucocytosis in response to the presence of bacteria. In the peritoneum. Phagocytosis in tuberculosis. CHEMOTAXIS. Botanical studies. Early studies of Leber. Early studies of Buchner. Methods. Theories of chemotaxis. Importance of surface tension.

CHAPTER XII.—PHAGOCYTOSIS, *Continued*. THE RELATION OF PHAGOCYTOSIS TO IMMUNITY 296

Opsonins and tropins. Metchnikoff's attempt to establish parallelism between phagocytosis and resistance. Work of his pupils. Metchnikoff's interpretations. Origin of bactericidal substances from leucocytes. "Macrocytase" and microcytase. Metchnikoff's interpretation of the Pfeiffer phenomenon. Origin of alexin. Leucocytic bactericidal substances. Their nature. Leucocytic ferments. Leuco-protease. Petterson's experiments. Leucocytic extract of Hiss.

CHAPTER XIII.—PHAGOCYTOSIS, *Continued*. FACTORS DETERMINING PHAGOCYTOSIS 311

Opsonins. Tropins. Metchnikoff's conception of stimulins. Work of Denys and his pupils. Other early observations. Work of Wright. Conception of opsonins definitely advanced. Analysis of opsonic action. Normal and immune opsonins. Neufeld's opinions. Bacteriotropins. Structure of opsonins. Specific absorption of opsonins. Heat stability of immune opsonins. Relation to other antibodies. Relation to alexin. Variations in leucocytes as a factor in opsonic measurements. Resistance to opsonic action on the part of bacteria. Relation to virulence.

CHAPTER XIV.—PHAGOCYTOSIS, *Continued*. OPSONIC INDEX AND VACCINE THERAPY 328

Wright's work on typhoid immunization. Development of technique for measuring phagocytic activity. The phagocytic index. Opsonic index. Dilution method. Simon and Lamar's method. Accuracy of opsonic index. Wright's work on the staphylococcus infections. Relation of opsonic index to clinical conditions. Negative phase. Summation of negative phase. Summation of positive phase. Clinical value of opsonic index estimations. Opsonins and tuberculosis. Treatment by auto-inoculation. The value of opsonic index determinations. The

value of vaccine therapy. Prophylaxis. Different types of infection and the logic of vaccine therapy in each type. The production and standardization of vaccines.

CHAPTER XV.—ANAPHYLAXIS. FUNDAMENTAL FACTS 358

The relation of immunity to hypersusceptibility. Various kinds of hypersusceptibility. Historical development of our knowledge of these phenomena. The work of Richet and others. The phenomenon of Arthus. The phenomenon of Theobald Smith. Experimental production of the anaphylactic state. Laws governing the condition as at first determined. Symptoms of experimental anaphylaxis in guinea pigs. Autopsy findings and causes of death. Changes in blood pressure. Changes in temperature. Leucopenia. Diminution of complement. Symptoms in rabbits and dogs. Anaphylactic antigen. Specificity of anaphylactic reaction. Quantitative relations. Variations depending upon method of administration. Anti-anaphylactic state. Prevention of anaphylaxis by drugs. Passive sensitization. Conditions governing its accomplishment. Quantitative studies of Doerr and Russ.

CHAPTER XVI.—ANAPHYLAXIS *Continued*. FURTHER DEVELOPMENT AND THEORETICAL CONSIDERATIONS 385

Theory of Gay and Southard. Besredka's theory. Gradual development of the antigen-antibody conception. Quantitative work. Identity of sensitizing and toxic substances. Idea of sessile receptors. Anaphylaxis and precipitins. The work of Vaughan. Diminution of alexin during anaphylactic shock. Toxic substances obtained by action of active serum. Friedberger's "anaphylatoxin." Obtained from precipitates. Obtained from bacteria. Is the mechanism of anaphylaxis intravascular or cellular? Precipitating and albuminolytins. Writer's opinion. THE MECHANISM OF ANTI-ANAPHYLAXIS. Nature of anaphylactic poison. Peptone shock. PHENOMENA CLOSELY RELATED TO ANAPHYLAXIS. Toxicity of normal serum. Toxin hypersusceptibility.

CHAPTER XVII.—ANAPHYLAXIS *Continued*. BACTERIAL ANAPHYLAXIS AND ITS BEARING ON PROBLEMS OF INFECTIOUS DISEASE 410

Early work on sensitization with bacterial protein. Technique for sensitizing with bacteria. Revision of our ideas of "endo-toxin." Vaughan's work on toxic protein split-products. Friedberger's anaphylatoxin. Methods of production. Quantitative proportions which must be observed. Time and temperature conditions. Bearing of this work upon our understanding of infectious disease. Friedberger's interpretation. Bacterial toxæmia. Is the bacterial antigen the matrix for the poison?

CHAPTER XVIII.—ANAPHYLAXIS *Continued*. THE CLINICAL SIGNIFICANCE OF ANAPHYLAXIS 426

Serum sickness. Accelerated reactions and immediate reactions. Methods of avoiding anaphylaxis in antitoxin injections. Anaphylaxis and bacterial vaccines. Asthma and hay fever. Sensitiveness to contact with certain animals. Possible anaphylactic reason for eclampsia. Sympathetic ophthalmia. Diagnostic reactions. Tuberculin reaction. Luetin reaction. Discussion of tuberculin reaction. Experimental anaphylaxis with tuberculin. Diagnostic use of anaphylaxis.

CHAPTER XIX.—THERAPEUTIC IMMUNIZATION IN MAN. THERAPEUTIC USE OF DIPHTHERIA ANTITOXIN 446

Statistical results. Amounts to be injected. Amount of antitoxin normally present in the human blood serum. PRACTICAL CONSIDERATIONS CONNECTED WITH DIPHTHERIA ANTITOXIN PRODUCTION AND STANDARDIZATION. Toxin production. L_0 and L_4 doses. Methods of determination. Production of antitoxin. Standardization of antitoxin,

U. S. Hygienic Laboratory method. Chemical concentration of antitoxic serum. ACTIVE IMMUNIZATION IN DIPHTHERIA WITH MIXTURES OF TOXIN AND ANTITOXIN. Behring's work. Use of the method. Results obtained. INTRACUTANEOUS METHOD OF DETERMINING TOXIN AND ANTITOXIN VALUES. Principles of the method. Uses. Application of the method to the determination of antitoxin in human beings. TETANUS ANTITOXIN AND ITS STANDARDIZATION. Determination of the unit. ANTITOXIN AGAINST SNAKE POISON. Calmette's work. Differences between cobra and rattlesnake poison. Production of antiserum. PASSIVE IMMUNIZATION IN DISEASES CAUSED BY BACTERIA WHICH DO NOT FORM SOLUBLE TOXINS. General consideration of principles involved. Difficulties. Serum treatment of epidemic meningitis. Work of Kolle and Wassermann. Experiments of Jochmann. Flexner and Jobling's experiments. Results. Present methods. Streptococcus antiserum. Differences between various races of streptococci. Marmorek's serum. Work of Aronson, Tavel, Van de Velde and others. Probable manner of action. Serum treatment in pneumonia. Neufeld's work. Recent experiments and methods of Cole. Serum treatment of typhoid fever. Earlier experiments. Attempts to produce anti-endotoxin. Principles involved. Immunization with trypsin digested bacteria. Immunization with sensitized bacteria. Prospects of success. Serum treatment of plague. Yersin's attempts. Kolle and Martini's serum. Work of British Plague Commission. Lustig's serum. General results obtained. FACTS CONCERNING ACTIVE PROPHYLACTIC IMMUNIZATION IN MAN. General principles. Typhoid vaccination. Earlier history. Work of Wright, Kolle, and others. Russell's report of vaccination in the United States army. Statistics. Work of Metchnikoff and Besredka. Prophylactic immunization against cholera. Methods. Results. Plague vaccination. Difficulties. Methods. Results. Smallpox vaccination. Rabies. Principles and methods of application.

| | |
|--|-----|
| CHAPTER XX.—ABDERHALDEN'S WORK ON PROTECTIVE FERMENTS. MEIO-STAGMIN REACTION | 493 |
|--|-----|

| | |
|---|-----|
| CHAPTER XXI.—COLLOIDS, by Professor Stewart W. Young, Stanford University, California | 499 |
|---|-----|

Introduction. Definition. Reversible and irreversible colloids. Stability of colloidal systems. Physical properties of colloids. Form and size. Osmotic pressure. Rate of settlement. Brownian movement. Electrical properties of colloids. Surface tension. Chemical properties of colloids. Flocculation of colloids by electrolytes. Salts and acid electrolytes. Influence of concentration. Difference in sensitiveness to electrolytes. Explanation of phenomenon. The "zone-phenomenon." Mutual reactions of colloids. Mutual flocculation. Protective action. Theories of interaction. The preparation of colloid solutions. Applications to biology. Living tissues as colloids. Agglutination of bacteria. Analogy to colloid phenomenon. Electrical charge carried by bacteria. Sensitiveness to light. Danysz phenomena. Conclusions.

INFECTION AND RESISTANCE

CHAPTER I

INFECTION AND THE PROBLEM OF VIRULENCE

THE early history of our knowledge of infectious disease is that of fermentation. It was a philosopher, Robert Boyle, writing in the 17th century, who prophesied that the problem of infectious disease would be solved by him who elucidated the nature of fermentation. His prediction was fulfilled 200 years later by the train of investigations begun by Cagniard-Latour and by Schwann, and carried to a brilliant culmination by Pasteur. It was the discovery of the living nature of ferments and the specific nature of the various micro-organisms which caused the several forms of fermentation, and especially of putrefaction, which made possible rational investigations in the field of infectious disease and led by analogy, first to logical speculation—then to actual experimental proof of the etiological relationship between the minute forms of life and the communicable diseases.

It is not much more than 50 years since Pollender described the anthrax bacillus in the blood and spleens of animals dead of this disease. In this short period the large number of maladies of animals and human beings caused by micro-organisms belonging both to the varieties spoken of as bacteria and to those classified as protozoa has necessitated the segregation of this branch of knowledge into a separate chapter.

The period of etiological investigation is now approaching its maturity. The causative agents of most of the more common infectious diseases have been discovered, and the biology of many of the pathogenic micro-organisms has been thoroughly studied both in their artificial cultures and in the infected animal body. In spite of a considerable accumulation of facts, however, the science of immunity, that is, the study of the defensive powers of the living animal body against infection, is still in its infancy, and the practical therapeutic successes based on this science are disappointingly out of proportion to the really large amount of detailed knowledge of cellular and serum reactions at our disposal.

The study of putrefaction and of fermentation—though furnishing the basic analogy from which the first impulse was obtained—

presented after all a problem infinitely more simple than that of the infection of living tissues with bacteria. For, given any organic material containing suitable nutritive constituents, with favorable environmental conditions of moisture and temperature, and spontaneously or experimentally inoculated with germs of a proper species, and the phenomena which ensued were merely those of bacterial growth, in which an active part was played by the bacteria only, the dead organic materials serving simply as a passive menstruum for these activities.

During the earlier days of the development of bacteriology, therefore, when the attention of investigators was concentrated primarily upon the discovery of the specific causal agents of various infectious diseases, it seemed that the simple bringing together of pathogenic germ and susceptible subject should suffice for the accomplishment of an infection. We have learned, however, that the process is much more involved, and that, fortunately for the survival of the higher animals and man, the conditions which determine infection are intimately dependent upon a variety of secondary modifying factors.

Throughout nature bacteria are abundant, and the environment of man and animals, the outer integuments of skin and hair, and the mucous membranes of the conjunctivæ, the intestinal and respiratory tracts, are constantly inhabited by a thriving bacterial flora. The distribution of certain species in definite localities is often sufficiently constant to be regarded as a normal condition. Thus the *Bacillus xerosis* is a characteristic inhabitant of the conjunctiva, certain cocci and spirilla are always present in the mouth and pharynx, as is Döderlein's bacillus in the vagina. The fact that bacilli of the colon group are invariably present in the bowels of animals and man from the first few days or hours after birth has even been interpreted by some investigators as a physiologically beneficial condition. In the course of ordinary existence, therefore, and much more so during the course of accidental exposure to individuals in whom infection is present, the bodies of the higher animals are in intimate contact, not only with ordinarily harmless bacteria (saprophytes), but also with many varieties of the micro-organisms spoken of as "*pathogenic*" or disease-producing. Perfectly normal individuals have, then, on occasion, been found to harbor diphtheria bacilli in nose and pharynx, meningococci have been found in similar localities, and tetanus bacilli, the bacillus of malignant edema, the Welch bacillus, and other distinctly pathogenic germs have been isolated from the intestinal contents of individuals who showed no evidence of disease. In fact, the problem of the so-called bacillus carriers—persons who, though themselves apparently well for the time being, harbor within their bodies and distribute to their environment bacteria capable of causing disease in others—is, as we shall

see, now recognized as one of the most important difficulties of sanitary prophylaxis. In the case of typhoid fever this is particularly true, for it is now well known that a perfectly healthy individual may harbor typhoid bacilli in the gall-bladder for years and constitute, through all this time, a constant focus of danger to the public health.

The accomplishment of an infection, then, is not determined merely by the fact that a micro-organism of a pathogenic species finds lodgment in or upon the body of a susceptible individual, but it is further necessary that the invading germ shall be capable of maintaining itself, multiplying and functioning within the new environment. An infection, then, or an infectious disease, is the product of the two factors, invading germ and invaded subject, each factor itself influenced by a number of secondary modifying circumstances, and both influenced materially by such fortuitous conditions as the number or dose of the infecting bacteria, their path of entrance into the body, and the environmental conditions under which the struggle is maintained.

We have in truth, then, a battle of two opposed forces, the result of which is infectious disease. And it is the systematic analysis of these forces in their variable conditions, and the laws which govern them, which constitutes the science of immunity. It is the initial skirmish between the two which determines whether or not a foothold shall be gained upon the body of the subject and an infection thus established, and it is the balance between them which decides the eventual outcome of recovery or death. And though it is unfortunately true that much of the knowledge gained by such studies has yielded no direct therapeutic results, the facts that have been revealed are fundamental to the pathology of infectious disease and as essential to the clinical understanding of these maladies as is the knowledge of the mechanism of the circulation, the chemistry of metabolism, or the structural changes of the tissues to the comprehension of other pathological conditions.

And from this point of view the study of infectious diseases can be made an eminently logical one, in that, knowing the criteria which govern the infection of a human being with a given germ, knowing the probable path of entrance, manner of distribution, and biological activities of the micro-organism, and the peculiarities of the mechanism of resistance set in motion in the body by this particular infection, definite clinical deductions can often be made.

One of the most fundamental facts, immediately apparent on considering the problems of infection, is the phenomenon that among the innumerable varieties of bacteria and protozoa present in nature there is a very limited group which is capable of becoming parasitic upon the body of higher animals, and among these a still smaller proportion which is capable of being "pathogenic" or causing disease. We have used the terms pathogenic and non-pathogenic as

practically synonymous respectively with "parasitic" and "saprophytic." But, as we shall see, although as a rule a micro-organism must be parasitic to possess pathogenic powers, some of the true saprophytes or so-called half-saprophytes may be pathogenic under certain conditions, and the terms do not cover each other absolutely.

It is reasonable to suppose that all micro-organisms were originally in the condition which we designate by the term "saprophytic." By this term we imply that these germs maintain themselves only upon dead organic matter and do not thrive in or upon the living animal tissues. The class of saprophytes is widely distributed and constitutes, of course, the most important group of bacteria in nature, since upon the activities of these germs depends the unlocking of nitrogen and carbon from the organic complexes in the dead bodies and waste products of animals and plants. Such bacteria if strictly saprophytic, that is, entirely unable to maintain themselves upon living tissues, have little importance as producers of disease, or, expressed in technical terms, have little "pathogenicity." Nevertheless, there are cases in which strict saprophytes may cause disease by lodging upon and growing in animal tissues which have been killed by other causes, so-called necrotic areas; and these, still being in relation with the body as a whole through the blood and lymph channels, furnish an area of saprophytic growth from which products of putrefaction or even bacterial poisons may be absorbed. While, as a rule, the disease following the invasion of necrotic tissue—such as gangrenous amputation stumps, old unhealed sinuses, diabetically gangrenous areas, etc., may be caused by a large variety of saprophytic bacteria, there are a few very important and specifically pathogenic bacteria which are, strictly speaking, saprophytes. Thus the form of meat poisoning caused by the *Bacillus botulinus* is due entirely to the poison formed by this bacillus outside of the body within the substance of the dead foodstuff, and disease ensues as the result of subsequent ingestion of this poison with the food. In the same way the tetanus bacillus and, less strictly speaking, the diphtheria bacillus, at least in its ordinary mode of attack, are rather closer to the class of saprophytes than to that of the parasites, since neither of these bacteria, under usual circumstances, invades the substance of the tissues beyond the point of initial lodgment, causing disease only by the production of specific poisons, a condition known as "toxemia" or intoxication. The tetanus bacillus, moreover, is not usually capable of maintaining itself and multiplying even at the point of initial lodgment unless the tissues have been injured by trauma or irritated by the presence of foreign bodies. Bacteria of such characteristics, therefore, though pathogenic—that is, incitant of disease—remain nevertheless essentially saprophytes living upon the dead animal tissues not invading the living cells or body fluids. It is true that investi-

gations of Frosch¹ have shown that diphtheria bacilli may often be found in blood and organs of diphtheritic patients, and tetanus bacilli have occasionally been found in the spleen. However, such distribution is not necessary for the production of disease by these bacteria, and the essential point remains that they may cause violent, often fatal, disease without truly departing from their saprophytic mode of life upon dead tissues. Between the saprophytes and the true parasites or invaders of living tissue many transitions occur, and the condition of parasitism is probably a form of specific adaptation.

How such transition may be biologically developed is probably well illustrated by the investigations of Italian bacteriologists upon tetanus bacilli.² Tarozzi³ inoculated guinea pigs and rabbits with tetanus spores subcutaneously and found that these spores were rapidly transported to the liver, spleen, and kidneys, where they could maintain a latent existence for as long as 51 days. If during this period trauma or any injury of the organs was practiced which led to the formation of necrotic tissue the spores would develop upon this basis and cause acute or chronic tetanus. Canfora,⁴ continuing these studies, likewise found that tetanus spores inoculated under the skin are rapidly distributed throughout the circulation. If no trauma has taken place at the point of inoculation the locally lodged spores may be rapidly destroyed, probably by phagocytosis. In the circulation they appear to be less rapidly eliminated and may be present for from ten to thirteen days. If, during this period, there is produced a small wound, blood clot, or necrotic area in the body—this may serve as a focus for development and tetanus may ensue. After ten or more days the spores disappear from the blood, but may then take up a latent existence in some of the organs—as stated by Tarozzi. Apart from their importance as constituting a sort of transitional condition between pure saprophytism and parasitism, these investigations would seem to have much bearing upon the so-called cases of “cryptogenic tetanus.”

True infection, that is, the invasion of one species by individuals of another, and the ability of the latter to multiply and functionate within the cell complexes of the former, is a process quite out of keeping with the ordinary plans of nature, throughout which there seems to be a distinct opposition to the colonization and functionation of one living being within the living substance of another. Thus, as Bail⁵ has pointed out, a mass of frogs' eggs will remain

¹ Frosch. *Zeitschr. f. Hyg.*, Vol. 13, 1893.

² Belfanti, quoted from Canfora, *Centralblt. f. Bact.*, I. Orig. Vol. 45, 1908.

³ Tarozzi. *Centralblt. f. Bact.*, Orig. Vol. 38, 1905.

⁴ Canfora. *Centralblt. f. Bact.*, Orig. Vol. 45, 1908.

⁵ Bail. “Das Problem der Bakt. Infection.” Klinkhardt, Leipzig, 1911.

entirely uninvaded while alive, though the water surrounding it may swarm with bacteria of many varieties, but when by some accident such a mass of eggs ceases to live, it immediately falls prey to bacterial infection. The same point is illustrated by the rapidity with which intestinal bacteria will spread throughout the body after death, when during life they have remained confined to the lumen of the intestine, or, at most, get into the portal circulation, to be destroyed in the liver. By the living cell, therefore, an opposition is offered to invasion by bacteria, a vital function which Bail has attempted to make clearer by formulating it as a law, referring to it as "Das Gesetz der Lebensundurchdringlichkeit." Upon what cell function this vital resistance to invasion depends is to a large extent a mystery. It would seem to rest in principle upon the fact that the invading cell meets the invaded one under conditions peculiarly adapted to the activities of the latter, and is overcome before conditions suitable for its own activities have been established. The conditions here are not unlike those observed in the case of digestive enzymes, a comparison which becomes more than an illustrative analogy when we consider that apart from the mere mechanical disturbance created by the presence of bacteria as foreign bodies the struggle between invader and tissue is largely one of enzyme against enzyme. Thus, for instance, the gastric juice does not act upon the mucous membrane of the stomach during life—but after death, at autopsy, partial digestion of this membrane by the pepsin is often seen.

Whenever this vital resistance or opposition is overcome, and micro-organisms enter the tissues or cells, an abnormal process is taking place, and this process is, strictly defined, infection. Nevertheless, it is by no means necessary that such infection should always be accompanied by manifestations of disease. It is true that, in most cases, the natural resistance is such that a struggle ensues by which the invader is destroyed or thrown off, or in which the invaded subject is functionally injured or even killed, and the accompanying evidences of such a struggle constitute what we know as infectious disease. But there are special cases, cases of adaptation, biologically speaking, in which neither invader nor host is seriously harmed.⁶ In the field of protozoölogy, especially, there are many examples of true parasites, that is, invaders truly maintaining their metabolism at the expense of the tissues and body substances of the host, which do not arouse reactions sufficiently vigorous to be termed "disease." Thus the *Trypanosoma Lewisi* may be found in the blood of rats⁷ without noticeably affecting the health of the animals, and other protozoa have similarly been found in organs and blood stream of a number of other apparently healthy animals. Although such conditions have been frequently spoken of as "infection

⁶ See also Bail, *loc. cit.*

⁷ Doflein. "Die Protozoen als Krankheitserreger."

without infectious disease," the distinction is probably one of degree only—there being some reaction on the part of the host even in the mildest cases, if only in the weakening by withdrawal of body substance, which distinguishes the infected from the uninfected animal. In other cases there may even be advantage to the host, following the infection, to the detriment of the invading micro-organism, a phenomenon most clearly illustrated by the invasion of the root hairs of leguminous plants by the Nitrogen-fixing "root-tubercle" bacilli, a condition in which, as Fischer says, the plant may be regarded as parasitic upon the bacteria.

The actual harm resulting from the infection must, to a large extent, depend upon the degree of adaptation to the new conditions of life possible on the part both of the invader and of the host. If the invader can acquire resistance to the defensive properties of the host, and the latter can be similarly adapted to the harmful effects of the invader, a prolonged condition of infection might ensue, a sort of truce without manifestations of the disease. Although this is conceivable, such mutual adaptation is probably very rare in human disease.

In cases of so-called chronic septicemia in which bacteria may be again and again isolated by blood culture from the circulation it is more than likely that the organisms are constantly present, not because they multiply or maintain themselves within the circulation, but rather because they are being continuously discharged into the blood from an established focus in the tissues—as, for instance, on a heart valve. We have examined the serum of patients with subacute and chronic septicemia (endocarditis), and often found powerful opsonic action against the invading germs even when the patient's own serum and leukocytes were used in the tests, evidence that the bacteria were probably being successfully disposed of after they had gained entrance into the blood stream. In rabbits, too, in our experience and in that of Miss Gilbert of this laboratory, it would seem that protracted septicemia is present only when secondary foci have been established from which the bacteria are constantly being discharged into the blood. This we believe is rather the rule and the establishment of a balance within the blood stream an exception. When bacteria *do* succeed in withstanding successfully the opposing forces active within the circulating blood their rapid accumulation, the collapse of the defensive mechanism, and death of the patient are probably the most common course.

The point of view which we have expressed in the preceding paragraph has been impressed upon us with particular insistence by the observation of certain cases of bacteriemia following infections of the middle ear, mastoid processes, and thromboses of adjacent veins. In such cases it appears that the blood may be flooded with bacteria which, nevertheless, disappear after the focus of infection has been

removed. We have recently had the opportunity to observe this case of septicemia caused by *Streptococcus mucosus*, in which blood culture plates showed very numerous colonies, and in which recovery followed promptly upon complete excision of the thrombosed vein. It would seem to us, therefore, that bacteriemia offers a rather better prognosis than was formerly supposed, at least in cases in which the focus is surgically accessible.

The same principle is illustrated in the ordinary clinical course of typhoid fever in the human being. Here the disease begins as bacteriemia. Very rapidly, usually within two weeks, the bacteria disappear from the blood stream and a high serum immunity is established in the patient. Nevertheless, the bacteria remain active, growing within definite foci in the tissues, where they are to a certain extent protected or inaccessible to the defensive powers so successfully active in the blood stream. At any rate the patient remains diseased and the bacteria can be isolated from the spleen, gall-bladder, and intestines at a stage when they are no longer present in the blood stream, and during which a measurement of the bactericidal and opsonic powers of the patient will reveal a serum immunity much higher than normal. Just why the organisms are protected from these influences in the tissues we do not know.

On the other hand, it is nevertheless true that a certain amount of actual adaptation between the bacteria and the body may take place and contribute to the chronicity of an infection. This seems to be shown especially by the experiments of Walker and others, which are referred to in other places, in which it was found that bacteria grown on immune sera gain a certain amount of resistance against the injurious properties of these substances, and evidence more directly bearing upon the question is furnished by the studies on the typhoid carrier state in rabbits made by Chirolanza,⁸ Blackstein,⁹ Johnston,¹⁰ and recently by Gay and Claypole.¹¹ The latter named writers found that they could regularly produce the typhoid carrier state in these animals if they first cultivated the typhoid bacilli upon a medium containing defibrinated rabbits' blood. Even in these cases, however, it is not at all improbable that the typhoid bacilli establish a permanent focus from which they are discharged into the blood stream.

An infectious disease, therefore, may be interpreted as the result of parasitism in which no such mutual adaptation has taken place, and in which the invasion of the host by the micro-organism is marked by a struggle, the local and systemic manifestations of which constitute the disease. The disease is an evidence of conflict

⁸ Chirolanza. *Ztschr. f. Hyg.*, Vol. 62, 1909.

⁹ Blackstein. *Bull. Johns Hop. Hosp.*, 1891.

¹⁰ Johnston. *Journ. Med. Res.*, 27, 1912.

¹¹ Gay and Claypole. *Arch. of Int. Med.*, 12, 1913.

tween the two forces, mild and locally limited if the protective powers far outweigh the invasive powers of the micro-organisms, violent if the balance is reversed. This conception is probably a correct one in the case of the large majority of diseases—those in which invasion is accompanied by more or less rapid and violent inflammatory and other reactions. In diseases like leprosy, tuberculosis, and a few others of the more chronic infections it is also possible that extensive invasion of the body depends, not so much upon the active invasive powers of the micro-organism, powers which we will attempt to analyze presently, but rather upon the fact that for reasons of insolubility and lack of irritating properties on the part of the invader no reaction is set up at first, and the invasion, though progressive, elicits no violent symptoms and no energetic opposition. The invader therefore progresses unopposed, becoming an incitant of disturbed bodily functions to the degree of actual disease only when it has gained a foothold in some organ and begun to proliferate, or has multiplied in such numbers that the cumulative effect of its toxic powers becomes manifest.

Such a conception would assign the slow and gradual but progressively invasive powers of such diseases as tuberculosis, leprosy, and syphilis in which systemic symptoms are manifest only after the disease has gained an extensive foothold, to the lack of acute physiological reaction resulting from the presence of the invading micro-organism. In the case of such infections as those caused by some of the yeasts or blastomyces we have seen foci of blastomycotic lodgment in the kidney and other organs surrounding which there was neither an accumulation of mobile cells—(leukocytes or lymphocytes)—nor any evidence of cloudy swelling or other injury, by poisons, of adjacent parenchyma cells. Here, as in tuberculosis or leprosy, the reaction induced by the presence of the micro-organisms is slow and gradual—expressed in an eventual fixed tissue-cell reaction and giant-cell formation—similar to that induced by insoluble foreign bodies. And it may well be that the progressive ability to multiply without arousing the invaded body to rapid and powerful reaction may account for the prolonged period of apparent well-being in the early stages of such infections and permit the invaders to pervade the body so extensively.

This point of view has been, we believe, most clearly expressed by Theobald Smith.¹² Bacteria may lack invasive or pathogenic properties and be, therefore, immediately destroyed after gaining entrance to the host. They may be powerfully invasive and because of lack of adaptation arouse a violent defensive reaction on the part of the host. "There is another type of parasite," Smith says, "which may dispense largely with both offensive and defensive processes. We can conceive of this type as exerting a metabolic activity approx-

¹² Theobald Smith. *Journ. of A. M. A.*, May, 1913, Vol. 60.

limiting so closely to that of the host that the latter reacts but slightly and then only after a long period of stimulation." Into this class he places the syphilis spirochæta and, in a somewhat modified sense, the tubercle bacillus.

We have seen, then, that a micro-organism may be pathogenic and still be saprophytic in its mode of life. In order that this can occur, however, it is necessary that it should possess the power of producing at the place of lodgment a poison or toxin which can be absorbed and cause disease. The condition which ensues is not, properly speaking, an infection, but rather a "*toxemia*," differing from the toxemias resulting from the ingestion of drugs or other poisons only in so far as the toxins are manufactured at some point of bacterial lodgment within the body of the victim. Typical tetanus and diphtheria, for instance, can be produced as readily by ingestion of the bacteria-free culture filtrates as by inoculation with the bacteria themselves. And although these bacteria may, on occasion, become invasive and thereby satisfy the criteria of true infection, this is not necessary for their pathogenicity.

In the large majority of bacterial diseases, however, it is necessary that the germs shall be capable of producing a true infection before they can become pathogenic, and it is our task therefore to attempt to analyze those bacterial attributes upon which the invasive power or virulence may be said to depend.

In the realm of infectious micro-organisms a wide range of cultural variations is encountered which indicates that some of these germs have adapted themselves very closely to the specific environmental conditions found in the living animal body, while others can take up with ease and under the simplest cultural conditions a purely saprophytic existence.

Many pathogenic micro-organisms have so far defied all attempts at cultivation in artificial media. These we cannot use for examples since it may well be that the failure of attempts in many of them may hinge upon such simple alterations of method as the exclusion of oxygen, the addition of fresh tissue, or the supplying of amino-acids, which have made possible the cultivation of the spirochæta pallida and the leprosy bacillus. But among those which we can cultivate there are many which require for successful cultivation the production of artificial conditions simulating closely those obtaining in the living body. Thus malarial plasmodia can be made to multiply only if furnished with uninjured human red blood cells, within which they can develop. The gonococcus requires, in its first cultures outside the body, a medium containing human protein; and the hemophile bacteria, among them the influenza bacillus, require hemoglobin. Other organisms like pneumococci, many streptococci, diphtheria bacilli, and many others, though easily grown on artificial media, are still fastidious in their requirements and develop

sparsely or not at all unless definite conditions of nutrient materials, temperature, reaction, and osmotic pressure are observed. On the other hand, typhoid, anthrax, and dysentery bacilli, staphylococci and numerous other pathogenic germs grow easily and luxuriantly on the simplest laboratory media and within a wide range of environmental variations.

Biologically considered, we could arrange the scale of adaptation to parasitic conditions on this basis and it would seem, *a priori*, that those bacteria which had thus adapted themselves most closely to the living body should be the most infectious. There is not, however, such parallelism, since many of the most powerfully invasive or virulent germs, for instance, the anthrax bacillus, have retained their capacity for saprophytic life to the fullest extent. It is more logical, therefore, to classify parasites, not according to their ability to revert to saprophytic conditions, but rather, as Bail¹³ has done it, on the basis of their relative powers of invading the living body. His classification, of course, implies that the position of each micro-organism in this scale must be determined with reference to a given animal species, since a germ which is highly infectious ("parasitic" in Bail's sense) for one species may be a "half-parasite" or even a pure saprophyte for another.

Briefly reviewed, his classification is as follows:

I. *Pure Saprophytes*.—(Necroparasites, superficial parasites, or external parasites.)

Micro-organisms which under no circumstances can be made to develop within the living tissues of a given animal. This does not exclude their pathogenicity for this animal, since, like the diphtheria or tetanus bacillus, they may develop and produce toxins on the basis of a localized area of dead tissues.

II. *Pure Parasites*.—Organisms like the anthrax bacillus or the bacilli of the hemorrhagic septicemia group which, implanted in small quantity in an animal, will rapidly gain a foothold, thrive, and spread throughout the body.

III. *Half parasites*, organisms which may be infectious if introduced into the animal body, but, not possessing this invasive power to the same degree as the preceding class, require the inoculation of considerable quantities, often a special mode or path of inoculation, or even possibly a preliminary reduction of the local and general resistance of the infected individual in order that they may multiply and become generalized. This class includes the large majority of the bacteria pathogenic for man.

This property of invasive power is spoken of as *virulence* in contradistinction to *toxicity*—the latter implying merely the ability to produce poisons, and not necessarily being associated with the power to invade.

¹³ Bail. *Loc. cit.*

In order that a micro-organism may be a true parasite in Bail's sense—or invasive—for any given species of animal it must of course possess certain basic cultural attributes which enable it to grow in the environment furnished by the host. For instance, a micro-organism which does not grow at temperatures below 37.5° C. cannot very well become parasitic upon cold-blooded animals. An excellent illustration of this influence of body temperature upon the invasive powers of bacteria is furnished by the different races of acid-fast bacilli which invade the bodies of man and of birds. The avian tubercle bacillus, for instance, is non-pathogenic for man and in cultures will not develop at temperatures below 40° C., which is about the body temperature of most birds. The human tubercle bacillus, on the other hand, is non-pathogenic for birds and ceases to grow in artificial cultures when the temperature is raised above 40° to 41° C. This is merely one of a number of examples which might be cited to demonstrate the necessity of simple cultural adaptation, as it influences the property of virulence. Again, it is probable that in order to develop in the animal body it is necessary that a micro-organism shall be capable of developing without free oxygen. While this point is not definitely certain, it is not probable that any of the virulent bacteria can be strict aerobes. As a matter of experience none of the pathogenic bacteria at present known are absolute aerobes—though many of them grow better in artificial culture when oxygen is freely present than when it is absent.

Furthermore, the conditions encountered by bacteria as they enter the animal body will vary considerably according to the path by which they gain entrance. Organisms entering by the intestinal canal are subjected to conditions of acidity or alkalinity, the action of digestive juices, of bile, and to competition with other intestinal bacteria, forces to which many pathogenic germs will succumb, while others may survive there and thrive. Those entering into the tissues by way of the skin and mucous membrane, on the other hand, encounter an immediately mobilized protective mechanism which, successfully resisted by some of them, might easily and quickly dispose of small quantities of other bacteria more resistant to conditions in the bowel. It is but natural for this reason that the accomplishment of an infection by any given germ must depend to a great extent upon its gaining entrance to the body by the path best adapted to its peculiar requirements.

The mechanical protection afforded by the coverings of skin and mucous membranes is as a rule sufficient to prevent the penetration of any bacteria which by chance may have found lodgment upon them. In the case of the most usual pyogenic cocci and many bacilli such protection is probably absolute, and a distinct break of continuity, such as a bruise or a wound, even though this may be too small to attract attention, is necessary for successful infection. In

the case of a very limited number of diseases infection seems to take place even through the unbroken skin, and the method, often spoken of as the vaccination method of Kolle, employed in many instances when it is desired to produce experimental plague infection in rats or guinea pigs, consists in merely rubbing a small amount of cultural material into a shaven area of the skin. However, in this case, as well as in other instances where mere massage of bacteria into unbroken skin has led to successful inoculation, it is more than likely that success has depended upon either microscopic lesions or possibly the violent introduction of the organisms into the sebaceous glands, the sweat glands, or hair follicles. The defense of intact mucous membranes, however, is by no means impervious. While many organisms can be implanted upon mucous membranes with impunity, there are a number of others that can cause local inflammations upon these and can further pass through them into the deeper tissues and thence into the general system. Thus gonorrhea is ordinarily a disease of implantation upon a mucous membrane, and diphtheria bacilli and streptococci give rise to localized disease on the pharyngeal and nasal mucosæ, the latter not infrequently penetrating from the initial point of lodgment upon the mucosa into the deeper tissues and the circulation, causing a condition of "septicemia" or "bacteriemia." For the experimental determination of the penetrative power of organisms through mucous membranes the conjunctiva has been a favorite test object, and it has been shown that plague¹⁴ and glanders,¹⁵ as well as hydrophobia, may be transmitted by simple instillation of infectious material into the uninjured conjunctival sac. In the case of hydrophobia¹⁶ it is related that in Paris a young man contracted hydrophobia by rubbing his eyes with a finger contaminated with the saliva of a rabid dog. In the case of syphilis, though often claimed, there is no positive proof to show that infection may take place through the uninjured surfaces. It has been definitely shown, however, that tubercle bacilli¹⁷ may pass into the lymphatics through the intestinal mucosa without there being any traceable injuries on this membrane.

It may well be, however, that even without the existence of demonstrable morphological lesions penetrability by micro-organisms may presuppose local physiological or functional injury, such as congestion or catarrhal inflammation.

Thus it is seen that the mechanical obstacle to the entrance of micro-organisms offered by skin and mucous membranes, though important and not to be underestimated, is by no means a perfect safeguard.

¹⁴ Germ. Plague Com. *Arb. a. d. kais. Gesundheitsamte*, Vol. 16, 1899.

¹⁵ Conte. *Rev. vétérin.*, Vol. 18, 1893.

¹⁶ Galtier. *Compt. rend. de la soc. biol.*, 1890.

¹⁷ Bartel. *Wien. Klinikhandt*, 1906-1907.

However, it is only very definite species of micro-organisms which can cause disease at all when introduced into the body by these paths. For, although the rubbing of plague bacilli into the skin, or the inoculation of a cut surface with streptococcal or glanders bacilli, will rapidly lead to progressive infection, similar inoculation with the typhoid bacillus or the cholera spirillum would lead to no such result. And, though the swallowing of pus cocci, pneumococci, and a number of other micro-organisms would be entirely without effect, similar ingestion of the typhoid and cholera organism would usually result in typical infection.

The path of introduction, therefore, is an important consideration in determining whether or not a given micro-organism may give rise to disease. It is necessary that the manner of gaining entrance be suited to the cultural and other peculiarities of the germ in question. In the case of cholera, for instance, the spirillum which causes this disease is peculiarly susceptible to the deeper defences residing in the body fluids and cells, and cutaneous infection by the small numbers of bacteria likely to be introduced in this way would promptly be checked by these agencies. In the intestinal mucosa, however, the cholera spirillum finds conditions most favorable for rapid multiplication, and the disease is caused by the inflammation and destruction of the mucous and submucous tissues by the poisonous substances emanating from the large numbers of cholera spirilla which die and are disintegrated, as well as by the absorption of these poisons into the circulation. The bacteria themselves, however, never gain a permanent foothold within the blood or other organs. In the case of typhoid fever the conditions are somewhat similar, although here, during the earlier weeks of the disease, we have an actual penetration of the bacilli into the circulation. This, however, probably takes place only after intraintestinal proliferation has taken place, which then, on the injured mucosa, represents a dose out of all proportion great when compared with the quantities that would spontaneously come into contact with the external surface of the body.

This leads us to another important factor concerning the invading forces, in the determination of successful infection, namely, that of the *quantity introduced or the dosage*.

In order to cause infection, even when the bacteria are of the variety known to produce disease or "pathogenic," and are brought into contact with the body by a path suitable to their peculiar requirements, the initial quantity introduced must be sufficiently large to preclude complete annihilation by the first onslaught of the defensive powers of the body. It is plain, therefore, that in the case of bacteria weak in power to cause disease, given the subject of infection and his defences as a constant, the quantities to be introduced must be larger than in the case of micro-organisms of violent disease-

producing properties. The dosage necessary to cause infection, therefore, is in inverse proportion to that property of bacteria spoken of as their "virulence." Thus we measure the degree of the so-called virulence of bacteria by determining the smallest quantity, measured by dilution of platinum loops or by fractions of agar slant cultures (both very inexact methods), which will still cause infection and death in susceptible animals of a standard weight. In the case of micro-organisms of extreme virulence, such as the anthrax bacillus or bacilli of the hemorrhagic septicemia group, the inoculation of a very small number of bacteria may suffice to initiate infection. Indeed, it has been claimed for the anthrax bacillus that the injection of a single bacterium will produce fatal disease in a susceptible animal. The inverse relation existing between the degree of virulence and the number of bacteria inoculated is well illustrated by the experiments of Webb, Williams, and Barber,¹⁸ carried out upon white mice with anthrax, by the method of inoculation devised by Barber.¹⁹ This technique consists in picking up single organisms with a capillary pipette under microscopic control, from a very thin emulsion of bacteria and injecting directly from the pipette through a needle puncture in the skin. While requiring a considerable degree of skill, the method, when successful, permits an actual accurate count of injected bacteria instead of the merely approximate estimate which can be made by consecutive dilutions of thicker emulsions. In their experiments with anthrax in white mice Webb, Williams, and Barber found that the inoculation of a single thread of anthrax bacilli (3 to 6 individuals) taken directly from the blood of a dead animal (that is, in the most virulent condition) would regularly cause death, and it was impossible for this reason to immunize with such bacilli. On the other hand, if taken from 12-hour agar cultures of the same strain such small quantities would often fail to kill. The brief period of growth under artificial conditions had sufficiently lessened the virulence of the bacilli so that 2, 3, and more threads could be injected without harm. And after several generations of such cultivation as many as 27 and more threads could be inoculated with impunity.

Another example of the measurement of relative degrees of virulence, by a method more commonly employed, may be illustrated as follows: The problem in which this particular measurement was used consisted in the comparison of the virulence of two strains of pneumococcus, one (N_2) successively passed through white mice, the other (N_1) kept alive for several weeks on serum-agar. To accomplish this graded quantities of 18-hour broth cultures of the two

¹⁸ Webb, Williams, and Barber. *Jour. Med. Res.*, 1909, Vol. XV.

¹⁹ Barber. *Kansas Univ. Science Bulletin*, March, 1907.

strains were injected into mice of approximately the same weight as follows:²⁰

| N ₁ | Result | N ₂ | Result |
|----------------|--------------|----------------|--------------|
| 0.1 c. c. = | dead 24 hrs. | 0.1 c. c. = | dead 24 hrs. |
| 0.05 c. c. = | lives | 0.05 c. c. = | dead 24 hrs. |
| 0.02 c. c. = | lives | 0.02 c. c. = | dead 24 hrs. |
| 0.01 c. c. = | lives | 0.01 c. c. = | lives |

This example further illustrates another important fact in connection with the problem of infection—namely, that within the same species of bacteria different races of strains may exhibit widely varying degrees of virulence. This has been known since the days of Pasteur, and it is indeed of great importance in the immunization of animals that weakly virulent strains of a given micro-organism may be used to produce a gradual immunity against the same species of bacteria in their fully virulent condition. Though observed in almost all species of bacteria such variations are especially noticeable in the cases of streptococci and pneumococci—organisms in which no two strains may be alike in infectiousness, and in which the injection of some strains into susceptible animals may produce no result whatever, while other strains will kill if administered in the smallest measurable quantities. To a large extent these fluctuations of virulence appear to represent degrees of adaptation on the part of the bacteria to the conditions met with in the living body; and the ease with which such variations can often be artificially produced would seem to furnish another proof that the property of infectiousness is a biological attribute of relatively recent acquisition. For, although no general statement of absolute accuracy can be made, it is a fairly uniform rule that races of pathogenic bacteria gain in virulence as they are passed through successive animals of the same species, and lose in virulence as they are preserved upon media under conditions of artificial cultivation.

Further showing this ability to rapidly adapt themselves is the observation that passage through animals of a certain species will enhance the virulence for this species, but often reduce it for animals of another kind. Among the earliest observations on this point are those of Pasteur²¹ in his work on rabies. He found that the virus of hydrophobia when successively passed through rabbits gained in virulence until a degree of maximum infectiousness was attained

²⁰ For making such accurate measurements we have recently found very useful the Precision syringe described by Terry, *Jour. of Inf. Dis.*, Vol. 13, 1913.

²¹ Pasteur and Thuillier, *Compt. rend. de l'acad. des. sc.*, Vol. XII, 1883.

beyond which it could no longer be enhanced. After only three passages through monkeys, however, the virulence of this "virus fixe" for rabbits was reduced almost to extinction. His experience with swine plague was similar. Swine plague bacilli successively passed through rabbits and pigeons gained enormously in virulence for these animals respectively, but lost in virulence for hogs.

There are numerous methods by which the virulence of micro-organisms can be attenuated by laboratory manipulations, and since many of them are of great importance in the active immunization of animals we will reserve their detailed discussion until we come to consider the methods of immunization themselves. Suffice it to say in this place that most methods of attenuation consist in subjecting the bacteria, in artificial culture, to deleterious influences, either of unfavorably high temperature, exposure to light or harmful chemical agents, or allowing them to remain in prolonged contact with the products of their own metabolism by infrequent transplantation. As a rule the attenuation which inevitably follows any form of artificial cultivation in the case of bacteria like streptococci or pneumococci can be delayed by preserving them in media containing sera or tissues. In the case of the pneumococcus, for instance, one of the best methods of conserving virulence in storage is to keep them either in a soft rabbit-serum-agar mixture, as practiced by Wadsworth, or, better still, to store them within the spleen of a mouse dead of pneumococcus infection, as recommended by Neufeld. The mouse is autopsied and the spleen kept in the dark and cold in a desiccator, under sterile precautions. This, again, as well as the enhancement of virulence on passage through the same species of animal—or the reduction of virulence for one species by passage through another—shows that such fluctuations are dependent upon a very delicate biological adaptation.

It is interesting, moreover, to look upon this process of adaptation as a sort of immunization of the bacteria against the defensive powers of the host, a conception early suggested by Welch. For just as the animal body may become more resistant to the offensive weapons of the invaders, so it is reasonable to suppose that the bacterial body may gradually develop increased resistance to the defensive mechanism of the host. And this, if it occurs, would of course lead to an increase of its invasive power or virulence. The increase of virulence by passage through animals would alone lead us to suspect that such acquired resistance to destructive agents on the part of the bacteria might be responsible for the enhancement, but additional evidence pointing in this direction has been brought by experiments in which it was shown that bacteria cultivated in the serum of immune animals not only gained in resistance to destruction by the serum constituents, but at the same time were rendered more highly pathogenic. Experiments of this kind were carried out by

Sawtchenko,²² by Danysz,²³ and by Walker.²⁴ The results of Walker are especially instructive. He worked with a typhoid bacillus which he cultivated for a number of generations upon the serum of a typhoid-immune animal, and found that after such treatment the organism had gained in virulence and lost in agglutinability by immune serum, and that a larger amount of specific immune serum was necessary to protect animals against it than sufficed for protection against normal typhoid strains not thus cultivated. We will refer to these results more in detail in a later chapter, since the conception will be easier to grasp when we have considered more fully the mechanism of defence at the disposal of the animal body.

That this power of gaining resistance against deleterious influences on the part of bacteria is not confined to their resistance to the animal defences alone is well shown by the experiments of Danysz²⁵ upon the immunization of anthrax bacilli against arsenic. In inoculating series of 50 tubes containing arsenic dilutions (ranging from 1 to 10,000 to 1 to 200) with anthrax bacilli Danysz found that up to 1 to 5,000 the arsenic increased the growth of the bacilli; in concentrations higher than this growth was inhibited. By gradually progressive cultivation of the organisms in increasing concentrations of arsenic he finally succeeded in obtaining growth in solutions five times more concentrated than those in which they would develop at first.

It is intensely interesting also that Danysz found, both in the case of his serum-resistant and arsenic-resistant strains, that, as they became less sensitive to the deleterious effects of these agencies, they were altered morphologically in that they developed capsules. Similar in significance to this is the very important observation that certain strains of *spirochaeta pallida* may acquire resistance against salvarsan or "606."²⁶ These so-called arsenic-fast strains are apparently unaffected by the injection of this preparation into the patient.

The experiments of Danysz were probably the first to call attention to the possible relationship of bacterial capsule formation to virulence, and this particular phase of the subject has since then been extensively studied. It is a matter of common observation that micro-organisms like the pneumococcus, the anthrax bacillus, some streptococci, and a number of other germs which are capable of producing capsules under suitable conditions are most virulent in the capsulated stage. As the strains are passed through animals and their virulence increases their ability to form capsules becomes more

²² Sawtchenko. *Ann. Past.*, Vol. 11, 1897.

²³ Danysz. *Ann. Past.*, 14, 1900.

²⁴ Walker. *Jour. of Path. and Bact.*, Vol. 8, 1903.

²⁵ Danysz. *Loc. cit.*

²⁶ Oppenheim. *Wien. kl. Woch.*, 23, 1910, No. 37.

and more apparent—whereas the diminution of virulence which takes place on artificial media is accompanied by a gradual loss of capsule formation. Organisms like the Friedlander bacillus which retain their ability to form capsules almost indefinitely in artificial culture moreover do not lose their virulence to any great extent as long as this property is preserved. It is also well known that capsulated bacteria are peculiarly insusceptible to the ordinary agglutinating powers of specific immune sera. This has been noticed, not only in the case of heavily capsulated bacteria like those of the Friedlander group or the streptococcus mucosus, but, in the case of plague bacilli—where capsulation is usually present only in cultures taken directly from the animal body and cultivated at 37° C., Shibayama²⁷ has found a direct relation between non-agglutinability and a slimy condition of the cultures. Cultures kept at 5° to 8° C. in the ice-chest were easily agglutinable and lacked the slimy property. Cultures kept at 37.5° C. were slimy and thready in consistency and were not as easily agglutinated by the same immune serum. Porges²⁸ later showed that inagglutinable, capsulated bacteria can be made amenable to the agglutinating action of the serum—which we may assume to indicate vulnerability by the serum if the capsule is previously destroyed by heating at 80° C. for about 15 minutes in $\frac{1}{4}$ normal acid.

Against the cellular defences, the leukocytes, capsulated bacteria seem also to be more resistant than are the non-capsulated. This has been especially studied by Gruber and Futaki,²⁹ who find that a capsulated bacillus is rarely taken up by a phagocyte even when these cells are apparently normal and able to take up the uncapsulated organisms. They go so far as to claim that, in the case of anthrax in rabbits, the development or absence of a capsule determines whether or not infection can take place. The same conclusion is reached in similar studies by Preisz,³⁰ who does not believe that anthrax bacilli can ever cause infection unless they possess the power of forming capsules. All this experimental evidence points strongly toward a probable direct relationship between capsule formation and virulence, in the sense that a thickening of the ectoplasm may in some way protect the bacteria from the destructive forces aimed at them by the cells and fluids of the invaded body.

As a matter of fact, even when no distinct capsule is visible, it is nevertheless possible that ectoplasmic changes may take place. This phase of the subject has been thoroughly discussed by a number of writers, more especially by Eisenberg.³¹ It appears that many

²⁷ Shibayama. *Centralbl. f. Bact.*, Orig. Vols. 38, 1905, and 42, 1906.

²⁸ Porges. *Wien. klin. Woch.*, p. 691, 1905.

²⁹ Gruber and Futaki. *Munch. med. Woch.*, 6, 1906.

³⁰ Preisz. *Centralbl. f. Bakt.*, Vol. 49, 1909.

³¹ Eisenberg. *Centralbl. f. Bakt.*, I, 45, 1908, p. 638.

bacteria, in which true capsule formation has not been observed, may show swelling or enlargement under conditions in which their offensive activities in the infected animal body are called into play.³² Radziewsky³³ has noticed such swelling of *B. coli* in fatal guinea-pig infections, and spoken of it as "one of the characteristic signs of infectiousness." Kisskalt³⁴ has described the same thing in the case of streptococci, and Eisenberg interprets this as signifying an ectoplasmic hypertrophy comparable in principle to capsule formation. He looks upon the ectoplasmic zone as a protective layer, and calls attention to the observation of Liesenberg and Zopf³⁵ who showed that capsulated strains of *leukonostoc mesenteroides* will withstand 85° C., a temperature at which uncapsulated forms are rapidly killed.

There is a considerable amount of evidence, then, which seems to indicate that the development of a capsule is at least one important method by which the bacteria can protect themselves against the onslaught of the defences of the invaded animal body and in so doing become more virulent.

It is not likely, however, that this merely passive increase of the resistance to injury on the part of the bacteria accounts for the entire train of phenomena included in an enhancement of virulence. It has been suggested by a number of observers that definite active offensive characteristics distinguish the virulent from the avirulent bacteria, in that the former may secrete, within the living body, substances by which the destructive powers of serum and leukocytes are neutralized or held at bay. A very definite suggestion of such a possibility we find expressed in the now classical paper of Salmon and Smith³⁶ on hog cholera immunity, published in 1886. They say: ". . . the germs of such maladies are only able to multiply in the body of the individual attacked, because of a poisonous principle or substance which is produced during the multiplication of these germs."³⁷ Bouchard formulated such a theory in 1893 by speaking of the "*produits sécrétés par les microbes pathogéniques*," substances which he found in cultures of virulent bacteria, and which seemed to reinforce the invasive powers of the germs. Kruse³⁸ also within the same year developed a similar idea. He assumed that bacteria may secrete enzyme-like substances which paralyze the destructive properties of animal serum, and in this way gain the power to

³² These forms Bail has spoken of as "thierische Bazillen."

³³ Radziewsky. *Zeitschr. f. Hyg.*, Vol. 34.

³⁴ Kisskalt. Cited after Eisenberg, *loc. cit.*

³⁵ Liesenberg and Zopf. *Centralbl. f. Bakt.*, Vol. XII, 1892.

³⁶ Salmon and Smith. *Proc. Biol. Soc.*, Washington, D. C., III, 1884, 6, p. 29.

³⁷ A typewritten copy of this paper was kindly put at my disposal by Prof. Theobald Smith.

³⁸ Kruse. *Ziegler's Beiträge*, Vol. XII, 1893.

invade. As a matter of fact we have learned, since that time, that staphylococci may secrete soluble substances, "leukocidins," which injure white blood cells, and that many bacteria produce similar poisons, "hæmotoxins," which specifically injure red blood cells—thereby causing anæmia and reducing the resistance of the host. However, the correlation and further elaboration of these thoughts of Salmon and Smith, of Bouchard and of Kruse was left to Bail,³⁹ in what is known as his "aggressin theory." Bail maintains on the basis of careful experimentation that virulent bacteria can produce within the animal body substances which he calls "aggressins," upon which depend their invasive powers or virulence. These substances are secreted only under stress of the struggle against the unusual defences, are not demonstrable in test-tube cultures, and are in themselves, according to Bail, entirely non-toxic.

He obtains these aggressins by injecting virulent bacteria into the peritoneal cavity of a guinea pig and immediately after death removing the exudate. This he centrifugalizes, removes the bacteria and cells, and sterilizes the supernatant liquid by the addition of small quantities of chloroform. The action of the exudates in which aggressins have been produced by the bacteria is the following: (We take this tabulation from Bail's own paper on typhoid and cholera aggressins in the *Archiv für Hygiene*, Vol. 52, p. 342.)

1. Sublethal doses of typhoid bacilli or cholera spirilla become lethal when the aggressin is injected with them.

2. Lethal doses of bacilli which ordinarily would cause a slow infection only cause a rapid and severe infection when aggressins are added.

3. The addition of aggressin neutralizes the bacteria-destroying power of immune serum in the peritoneal cavity of a guinea pig.

4. The injection of aggressin alone produces subsequent immunity.

It is impossible to discuss with completeness the arguments advanced for and against the correctness of Bail's views until we have described in detail the mechanism of protection at the disposal of animals. But the main objection brought against this theory is that of Wassermann and Citron,⁴⁰ who claim that all these properties of the aggressive exudates can be explained by the fact that they contain extracts of the bacteria (endotoxins), which, injected with a sublethal dose of bacteria, merely enhance their action in the same way that this would have been accomplished by the injection of additional dead bacterial bodies. It will require much further work before this point is settled, and the problem is peculiarly involved and

³⁹ Bail. *Archiv f. Hyg.*, Vols. 52 and 53, 1905; *Folio serologica*, Vol. 7, 1911.

⁴⁰ Wassermann and Citron. *Deutsche med. Woch.*, Vol. 31, 28, 1905.

difficult. However, the recent work of Rosenow⁴¹ on pneumococci seems to bring some reënforcement to the ranks of those who maintain the existence of a special offensive substance at the command of virulent bacteria. Rosenow extracted pneumococci grown on serum broth and found that such extracts when made from virulent strains would protect avirulent strains from engulfment by phagocytes. The non-virulent strains left in these extracts for 24 hours became virulent. He believes, therefore, that the virulence of pneumococci depends largely upon the possession of these substances which he calls "virulins," and which in function at least are conceived as very similar to the "aggressins."

Recent results obtained by the writer⁴² with Dwyer seem to indicate that anaphylatoxins produced from the typhoid bacillus possess some of the properties claimed for his aggressin by Bail. It is not impossible that the "aggressins" obtained by him were of this nature.

Virulence, then, may be analyzed into two main attributes: one a purely passive property of resistance or self-preservation on the part of the bacteria, perhaps morphologically expressed in ectoplasmic hypertrophy and capsule formation; the other an actively offensive weapon in the form of substances of the nature of the "aggressins" of Bail or the "virulins" of Rosenow. The extent of our present knowledge of details does not warrant a statement of the case in more definite terms.

From the facts we have discussed in the preceding paragraphs it now becomes manifest that the elements which determine the nature of an infectious disease are twofold. On the one hand each variety of infectious germs possesses certain biological and chemical attributes which are specific and peculiar to itself; by these its predilection for path of entrance and mode of attack is determined, and upon these depends the nature of the reaction called forth in the animal body. On the other hand the degree of infection in each case, the severity of the reaction and the ultimate outcome are determined by the balance which is struck between the virulence of the entering germ and the protective mechanism opposed to it.

The specific properties of each micro-organism are the factors which account for the clinical uniformity (within definite limits) which is observed in the maladies produced in different individuals by the same species of bacteria. Thus a severe typhoid fever is, in essential characteristics, entirely similar to a mild case—since in both instances the path of entrance, through the intestine, is the same, the distribution of the germs after entrance differs only in degree, and the reactions, local and systemic, which are called forth

⁴¹ Rosenow. *Jour. of Inf. Dis.*, Vol. 4, 1907.

⁴² Zinsser and Dwyer. *Proc. Soc. Exp. Biol. and Med.*, Feb., 1914.

are alike. And cases of this disease in general differ as a class from the maladies caused by, let us say, the group of clinical conditions resulting from anthrax infection, where entrance is through the skin, and generalized infection of the blood ensues without definite or regular localization in any given organ. Again, a localized staphylococcus abscess will differ materially from an equally localized focus of tuberculosis, because the chemical constituents of these bacteria respectively call forth each a characteristic response on the part of the defensive mechanism.

Such specificity of the various micro-organisms may of course be due partly to their mode of attack and distribution, and partly, as we shall see, to the pharmacological action of the poisonous products given out by them.

That both factors contribute seems beyond doubt; but recent work, especially that of Friedberger, which is fully discussed in another place (see p. 413), seems to show that clinical differences depend much less than was formerly supposed upon specificity of the intracellular poisons, and much more upon distribution and localized accumulation of the germs, conditions which are determined rather by the mode and extent of invasion than by chemical differences of poison production. This problem, rather difficult to discuss on the limited basis of the facts so far outlined, will become clearer as we proceed, but we need only refer at present to the essential clinical uniformity of the various forms of septicemia, where organisms freely circulate in the blood—with often a focus of distribution on a heart valve—conditions in which it is rarely possible to determine the species of the responsible germ except by blood culture. Or, again, as Friedberger⁴³ points out, there is great similarity between the ordinary pneumococcus pneumonia and that caused by the Friedlander bacillus. In both cases the distribution and mode of attack of the bacteria are essentially the same, though the micro-organisms themselves are biologically very dissimilar.

One and the same micro-organism, on the other hand, may cause entirely different clinical conditions, and here the type of infection depends purely on the degree of invasion possible in the given case—that is, the balance between virulence and resistance. A germ may enter the body and cause an inflammatory reaction at the point of entrance, the process remaining purely localized. In such cases the defensive forces have been so efficient, the invasive properties of the germ so relatively weak, that progression beyond the point of entrance is prevented and the resultant disease takes the form merely of a localized abscess. This is the case when a healthy individual is infected with an attenuated organism or by one whose species' characteristics do not include a powerful invasive property. Thus streptococci, if entering the tissues of a normal subject in small

⁴³ Friedberger. *Deutsche med. Woch.*, No. 11, 1911.

numbers or in attenuated form, may produce a purely localized infection, and ordinarily non-pathogenic germs like proteus, subtilis, or colon bacilli may produce localized abscesses in weak and debilitated individuals, though implanted upon a healthy subject they would be rapidly disposed of without gaining even a preliminary foothold. Such tendency to localization is the common form of infection in the case of a number of germs. It is the most usual type of staphylococcus infection, for instance, in which the degree of virulence of the strains ordinarily met is such that the balance struck by them with the average defensive powers of man results in localization. However, the same micro-organism, enhanced in virulence, or gaining entrance in unusual numbers in a weakened individual, may rapidly spread from the point of inoculation, at first by contiguity, then by invasion of the blood and lymph channels, and become generalized.

When organisms become generalized and circulate in the blood the resulting condition is spoken of as septicemia or bacteriemia. This is the form of infection commonly caused by streptococci, bacilli of the hemorrhagic septicemia group, anthrax bacilli, and many others. It implies a powerful invasive property and always constitutes a condition of great gravity when persistent. We are learning of recent years, however, that in many infectious diseases formerly regarded as purely localized a temporary entrance of the bacteria into the circulation is a usual occurrence. Thus Fraenkel⁴⁴ has shown that lobar pneumonia is almost always accompanied during the acute stages of the disease by pneumococcus septicemia, and in typhoid fever we now know that the organisms circulate freely in the blood during the first two weeks of the disease, and often longer than this.

In these and other conditions the bacteria may be gradually destroyed and disappear from the blood stream as the immunity of the subject increases. In other cases the bacterial activities may be partially checked, the process becoming slower and more chronic. This is especially often the case when micro-organisms after entrance to the circulation have found a secondary lodgment upon a heart valve, from which a continuously renewed supply of bacteria can be given off to the blood. A special form of such "malignant endocarditis" caused by the *Streptococcus viridans* is particularly apt to take this chronic course.

The presence of bacteria in the blood is not, therefore, as formerly supposed, an invariably fatal condition.

Adami's recent work would indicate, moreover, that bacteria may normally enter the portal or even the general circulation from the intestine during health. This condition of "sub-infection," as he calls it, is more fully discussed on p. 234. That colon and other in-

⁴⁴ Fraenkel. *V. Leyden Festschr.*, 1902.

testinal bacteria may often penetrate into the portal circulation is indicated by the occasional occurrence of colon bacillus abscesses after trauma of the liver. In most septicemias, however, caused by virulent bacteria the invasion of the blood stream persists, rapid multiplication occurs and leads to death.

From the circulation the bacteria may gain lodgment in various organs and cause the formation of secondary abscesses. This condition is known as "pyemia," and may be caused by almost any bacteria which are capable of producing septicemia. Thus staphylococci, streptococci, or pneumococci may lodge in bones, joints, brain, or kidneys, in fact in any organ in which they can gain a foothold. However, there are evidences of distinct tissue predilections on the part of certain germs. Thus the virus of rabies and that of poliomyelitis, though to some extent universally distributed, seem especially to concentrate in the nervous system; cholera spirilla and dysentery bacilli appear to find conditions most favorable for development in the intestinal mucosa; amebic abscesses are most common in the liver; gonococcus infections when generalized find secondary localization with particular frequency on heart valves and joints; leprosy bacilli have a predilection for the nerve sheaths; and glanders bacilli injected into the peritoneum of a male guinea pig localize with such regularity in the testicles that the experiment has diagnostic value (Strauss test). Conversely it is only explicable on the assumption of such selective lodgment that tubercle bacilli, even though otherwise universally distributed through the body, will be absent from striped muscle tissue, and rare in the walls of the stomach. Such selection—as far as we can account for it at all, seems to depend upon the varying cultural conditions encountered by the germs in different organs.

On the other hand, localization may also be dependent upon accidental conditions such as trauma. Infections in which the entrance of bacteria is coincident with injury—as in the case, for instance, of compound fractures—will be able to spread throughout the injured region much more easily than they could enter the healthy tissue. In fact, it is well known that local tissue injury at the point of inoculation favors infection since it furnishes a rich substratum for growth in the form of dead cells or blood clot and interferes with the accomplishment of a normal protective reaction. In cases in which bacteria are circulating in the blood mechanical injury may create a focus of reduced resistance on which the invaders can gain a foothold. It is in this way perhaps that, among other things, we can explain tuberculosis of joints or bones which present a history of injury preceding the development of the infection—or the pleurisy and lobular pneumonias which have been known to ensue upon the fracture of a rib.

It is also possible that bacteria may be distributed in various

organs directly from the initial focus by embolism or by the massive invasion of a blood vessel. It is by such breaking into a vein that Weigert explains the generalization of miliary tuberculosis.

The inflammatory reaction which usually ensues at the point of entrance of bacteria is merely a result of the local struggle between invader and tissues, and the violence of this reaction is in a large measure an indication of the resistance of the infected subject. When, for instance, a streptococcus of moderate virulence gains lodgment in the skin of a healthy individual the rapid mobilization of leukocytic and other defences may prevent further invasion by the bacteria and lead to a struggle which is clinically evidenced by severe local symptoms. Did the virulence of the streptococci far overbalance the powers of resistance the local struggle might be reduced to a minimum, the infection progressing without any, or with but a slight local, reaction. The fact that pneumococci lodging in the human lung ordinarily cause lobar pneumonia is merely an evidence of a considerable degree of resistance to these germs on the part of the average human being. Pneumococci introduced into the pulmonary alveoli of very susceptible animals (rabbits) may pass directly through into the circulation, causing fatal septicemia without leading to a more than mild and temporary reaction in the lungs themselves. If, as in Wadsworth's⁴⁵ experiments, the rabbits are partially immunized—that is, their resistance increased before the pulmonary inoculation is carried out—a violent local reaction, analogous to lobar pneumonia, may follow, the severity of the reaction at the portal of entry being manifestly an evidence of more energetic opposition to further penetration of the bacteria.

The entrance of bacteria into the deeper tissues, and even the circulation, without any, or with but slight, local evidences of infection at the point of entrance is by no means rare. The innocent appearance of the site of the entrance of the bacteria in generalized streptococcus infection is a common surgical observation, and a streptococcus-infected wound of the hand or leg in a patient dying of septicemia may appear but slightly inflamed and edematous and incomparably milder in appearance than a staphylococcus boil with which the patient is walking about and suffering hardly any systemic disturbance.

Between the time of entrance of the bacteria into the body and the first appearance of symptoms of disease there is always a definite interval which is spoken of as "incubation time." This period is made up of two definite divisions—one the time necessary for growth, distribution, and accumulation of the bacteria, the other the time necessary for the action of the toxin or poison which may be secreted. The latter, the incubation time of the toxin, is a subject which is still unclear in many of its phases, and will be discussed

⁴⁵ Wadsworth. *Am. Jour. of the Med. Sc.*, Vol. 27, 1904.

in the following chapter (see p. 37). The former, however, is easily comprehended, in fact, is to be expected. For the small number of bacteria which gain entrance to the tissues in spontaneous infection is entirely inadequate in itself to produce symptoms. It is necessary that multiplication shall take place until the bacteria have accumulated in number sufficient to cause noticeable physiological disturbance. That the interval necessary for this must vary according to the number of bacteria originally introduced, the virulence of these, and the specific resistance of the patient goes without saying. Von Pirquet and Schick have suggested also that the incubation time may correspond roughly to the interval during which the subject is becoming "allergic" or hypersusceptible to the bacteria or virus. This will be discussed at greater length in the chapter on anaphylaxis.⁴⁶

But within the limits of the variations introduced by these factors the incubation time of each infectious disease—if spontaneously acquired—is sufficiently uniform to be characteristic. Thus the primary lesion in syphilis follows the inoculation after an interval of two to three weeks, rabies follows inoculation with street virus after about four to six weeks, the period being somewhat dependent on the location of the bite; typhoid fever takes about two weeks to develop; gonorrhea about five to seven days; small-pox about two weeks; yellow fever three to five days; and scarlet fever and diphtheria about two to six days. In general, it may be stated that within the limits observed for each particular infection the shorter the incubation time the more severe is the infection. Thus if tetanus follows inoculation with the tetanus bacillus within seven days the prognosis is far more grave than when the incubation time has occupied two or three weeks. And if localized and general symptoms follow rapidly (within twenty-four to forty-eight hours) after a streptococcus infection it is likely that the process is a very severe and virulent one.

⁴⁶ Von Pirquet u. Schick. *Wien. kl. Woch.*, 16, 1903, pp. 758 and 1244.

CHAPTER II

BACTERIAL POISONS

WHEN bacteria have gained a foothold anywhere within the animal body the local and general disturbances which follow, in all but the mildest and most trifling cases, are such that we cannot account for them solely on the basis of mechanical injury.

It may well be that the obstruction of capillaries and lymphatics and the pressure upon parenchyma cells, always incident to inflammatory reactions, contribute materially to local destruction, and thereby indirectly to systemic effects. However, even in diseases like anthrax, in which the body of the victim after death is found flooded throughout with masses of bacteria, these factors cannot fully explain the clinical manifestations. And such cases, indeed, are extreme examples, since, in the large majority of bacterial diseases, the illness resulting in the patient is severe out of all proportion to the extent of the tissue area invaded.

Moreover, all infections, if at all severe, whatever their nature or localization, give rise to fever, and this symptom alone, if carefully observed from hour to hour, may be sufficiently characteristic to indicate the specific micro-organism which is causing the illness. With this there occur alterations of the blood picture, either a numerical increase of white blood cells (leukocytosis) or a change in the relative proportions of the different kinds of leukocytes—or again an anemia caused by the destruction of red cells. There may also be degenerative changes in parenchyma cells of organs far removed from the actual site of bacterial lodgment. All these facts indicate very definitely that, apart from localized tissue destruction or purely mechanical interference with function by capillary obstruction or pressure, there is at the same time an absorption of poisonous substances emanating from the bacteria.

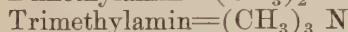
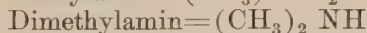
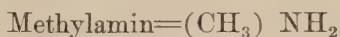
From the earliest days of logical investigation into the nature of infectious disease, as soon, in fact, as cultural methods had been introduced, bacteria were studied with the purpose of throwing light upon this phase of their activity. As a result of such investigations Selmi,¹ in 1885, described certain basic toxic substances which he obtained from putrefying human cadavers and for which he suggested the designation "*ptomain*" (from *πτῶμα* = dead body). These

¹ Selmi. Cited from Hammarsten, "Textbook of Physiol. Chem.," p. 16.

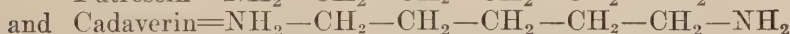
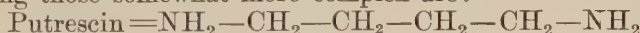
poisons were later more extensively studied by Brieger,² Gautier,³ Griffiths,⁴ and others, and it was at first surmised that the formation of such substances in the infected animal might be held responsible for the toxemic manifestations which accompany bacterial disease.⁵

This, as we shall see, is not the case. Ptomains are probably not formed in traceable quantity in the living tissues and are not in any way identical with the specific bacterial poisons which are responsible for the toxemia of infectious diseases. Nevertheless, they have some pathogenic significance, since they are invariably products of the proteolysis caused by bacteria and can give rise to illness when ingested with putrefying foodstuffs. It is important, therefore, that we discuss them briefly and consider their fundamental distinction from the true bacterial poisons.

Whenever dead organic material, meat, fish, vegetable refuse, etc., is left to itself under suitable conditions of moisture and temperature, putrefaction sets in. As a result of bacterial growth the protein is broken up and among the intermediate products of such proteolysis ptomains appear. Chemically^{6 7 8} these substances are basic nitrogenous compounds which may or may not contain oxygen. Because of their basic and often highly toxic properties they have been spoken of as "animal alkaloids." Many of them contain only C, H, and N, and are ammonia substitution products. (See Vaughan and Novy, *loc. cit.*, p. 248.) Thus some of the simpler ones are:



Among those somewhat more complex are:



Samuely classifies the ptomains according to their nitrogen contents as follows:

1. Those with one nitrogen atom ($\text{C}_8\text{H}_{11}\text{N}$) ($\text{C}_8\text{H}_{13}\text{N}$) ($\text{C}_{10}\text{H}_{15}\text{N}$)

2. Those with two nitrogen atoms such as putrescin ($\text{C}_4\text{H}_{12}\text{N}_2$) and cadaverin ($\text{C}_5\text{H}_{14}\text{N}_2$) and

² Brieger. "Die Ptomaine," Berlin, 1885; *Virchow's Archiv.*, Vols. 112 and 115; *Berl. klin. Woch.*, 1887, 1888.

³ Gautier. Cited after Pick, *Bull. de l'acad. de m d.*, 1886.

⁴ Griffiths. *Compt. Rend. de l'acad. des sc.*, Vol. 113.

⁵ For a historical outline of our knowledge of these poisons, as well as for a thorough treatment of their nature, see Vaughan and Novy, "Cellular Toxins."

⁶ For a discussion of the chemistry of the ptomains see Vaughan and Novy, "Cellular Toxins," Lea Bros., Philadelphia, 1902.

⁷ Also Samuely in Oppenheimer's "Handbuch der Biochemie," Vol. I, pp. 794 *et seq.*

⁸ See also Wells, "Chemical Pathology," Saunders, Phila., 1907.

3. Those with three nitrogen atoms such as methyl guanidin ($C_2H_7N_3$).

4. Finally there is an important group which contains oxygen, such as the substance sepsin ($C_5H_{14}N_2O_2$) obtained by Faust from putrefying yeast cells.

They are not in all cases protein cleavage products, since bodies of the cholin group, cholin, neurin, and muscarin, the two last named highly toxic, are lecithin derivatives, and Samuely points out that other lipid cleavage products, always present in decomposing tissues, may well contribute to ptomain production in the presence of a source of nitrogen. It is interesting to note also that the vegetable poison muscarin, isolated by Schmiedeberg from mushrooms, is chemically identical with a toxic base found by Brieger in decomposing fish.

The ptomains are not poisonous in every case. The chemically simpler ones like methylamin, di- and trimethylamin possess little or no toxicity. Others chemically more complex—like cadaverin and putrescin—may be capable merely of causing local necrosis, while sepsin, closely related to cadaverin in chemical constitution, but containing oxygen, is a powerful poison which acts violently upon the intestinal blood vessels, causing capillary dilatation, congestion, and diapedesis.⁹ The presence of oxygen seems indeed to be necessary for the development of strong toxicity (Brieger, Vaughan, and Novy). Again, the lecithin derivative, cholin, is but weakly toxic, while neurin is exceedingly poisonous. In putrefying mixtures these toxic bodies appear on or about the fifth or seventh day after putrefaction sets in, and disappear, by further cleavage, more or less rapidly, yielding less complex nitrogenous substances that are non-toxic.

With the limited knowledge regarding bacteria and infectious diseases at the disposal of the earlier investigators it was but natural that the discovery of ptomains in cultures of putrefactive bacteria aroused the suspicion that these bodies were responsible for the toxemia of infectious disease.

The search for poisonous substances in pure cultures of pathogenic bacteria was, therefore, assiduously taken up by Brieger and his pupils, and, in truth, ptomains were actually found as products of some of the disease-producing micro-organisms, just as they had been found in the mixed cultures involved in the putrefaction of meat. Thus cadaverin was found in cultures of the cholera spirillum, another nitrogenous poison, typhotoxin, in those of typhoid bacilli, and still another in tetanus cultures, all of them producing more or less severe illness when injected into animals.

In spite of this evidence, however, we have been forced to conclude that the ptomains cannot properly be held responsible for bac-

⁹ Meyer and Gottlieb. "Experim. Pharmacologie," 2d ed., p. 262.

terial toxemia as manifested in disease. In the first place it is doubtful whether ptomains, in noticeable quantity, are ever produced within the living infected body. Then, again, potent ptomains are produced in culture by many bacteria having absolutely no pathogenic power, while highly pathogenic bacteria may produce little or no ptomains. Ptomain production, moreover, is not specific, since the same ptomains may be produced by many different bacteria or mixtures of bacteria, provided the conditions of nutrient materials and temperature are favorable for growth. We cannot therefore account for bacterial toxemia, in which the poison produced by an individual species is characteristic and invariably the same, under varying cultural and environmental conditions, by the production of ptomains. And even when ptomains are produced in culture fluids by pathogenic bacteria their physiological action is usually quite different from that of the poisons produced by the same micro-organisms in the infected subject.

Briefly summarized, therefore, the ptomains are poisons elaborated by all bacteria that are capable of producing protein cleavage, if planted on suitable nutrient materials under conditions favoring growth. The matrix of these poisons is the protein nutriment; they are not products of intracellular metabolism specifically characteristic of the bacteria which produce them.

Their importance in the production of disease, therefore, is really an indirect one. They may cause disease if putrid meat or other material is ingested, and with it preformed ptomains, which may be taken in and further elaborated by continued putrefaction in the intestines. This form of meat poisoning, without bacteriological investigation, may be difficult to distinguish from such bacterial forms of meat poisoning as those caused by the Gärtner bacillus or the bacillus botulinus. Novy¹⁰ believes that true ptomain poisoning of this kind is rather less frequent than formerly supposed. However, in such cases as those of Vaughan, who isolated a poisonous ptomain "tyrotoxin" from cheese and milk, their importance seems reasonably certain. It is also probable that certain forms of auto-intoxication may be caused by the production in the intestinal canal of ptomains resulting from bacterial putrefaction incident to faulty digestive conditions. It is the antagonism to such intestinal putrefaction by the acid production of the bacillus *Bulgaricus* which is probably the basic cause of any favorable therapeutic effects which have attended the soured milk therapy of Metchnikoff. Again the growth of saprophytes in necrotic tissues such as gangrenous extremities in diabetes or amputation stumps, may lead to the formation of ptomains which, after absorption, can cause disease. In all such cases the process is one determined by the bacterial putrefaction of dead organic materials, and the absorbed

¹⁰ Novy in Osler's "Modern Medicine," Vol. 1, p. 223.

poisons are not true bacterial toxins, since they do not emanate specifically from the cell substance of the micro-organisms but rather represent incidental cleavage products of the nutrient materials. Therefore, also, the ptomains are unspecific—their formation a common attribute of a large variety of saprophytic organisms, their production, as to quantity and kind, primarily dependent upon the nature of the nutrient materials on which the bacteria are grown.

In contradistinction to the ptomains, the specific bacterial poisons, in the technical meaning of the term, are substances which are characteristic for each individual species of bacteria and truly the products of bacterial metabolism in that they emanate from the cell itself, either as a secretion or excretion during cell life, or as an inherent element of the cytoplasm liberated after death (or possibly as a cleavage product of the disintegrating bacterial protein).¹¹ They are dependent upon the nature of the culture medium only in so far as this favors or retards the normal development of the micro-organisms. While, therefore, a diphtheria bacillus undoubtedly produces the largest quantities of its specific poison on bouillon suitably prepared for this particular purpose, it will also, in smaller amount, produce qualitatively the same poison on all media on which its growth is free and uninhibited, even on a medium such as that of Uschinsky, which is entirely devoid of proteins. The *toxins* are, therefore, elements of intracellular metabolism, permanently or transiently constituent parts of the cell body.

A specific bacterial toxin was first obtained from the diphtheria bacillus by Roux and Yersin¹² in 1889. They discovered that if diphtheria bacilli were grown on veal broth and the cultures filtered through porcelain candles, after seven days at 37.5° C. the filtrates were highly toxic, producing the same symptoms and autopsy findings in rabbits, guinea pigs and birds which followed the injection of the living bacilli themselves. The poison was therefore a soluble product of the bacteria during the period of their vigorous growth, apparently given up by them to the culture fluid. Very soon after this, in 1891, Kitasato¹³ discovered a similar specific toxin in culture filtrates of the tetanus bacillus, and it was the hope of bacteriologists that analogous poisons could be determined for all pathogenic bacteria.

This hope, however, has been disappointed. It was soon found that cultures of cholera spirilla, typhoid bacilli, and many other germs did not yield toxic filtrates of this kind but that the poisons in these cases seemed to be firmly bound to the bacterial bodies dur-

¹¹ In connection with this read the discussion on anaphylaxis in chapter XVII, p. 413.

¹² Roux and Yersin. *Ann. de l'Inst. Pasteur*, Vol. 2, 1889.

¹³ Kitasato. *Zeitschr. f. Hyg.*, 1891, Vol. 10.

ing life, and given up to the surrounding media only after death and disintegration of the cells.

Pfeiffer¹⁴ was the first one to formulate this conception in his studies upon cholera poisons. He found that when cholera spirilla were grown upon broth and filtered after 6 or 7 days, the filtrate was but slightly toxic, but that, in this case, unlike the conditions prevailing in diphtheria and tetanus cultures, the residue of bacterial cell bodies, even after they had been killed by chloroform, thymol, or drying, were powerfully poisonous.

We have then two main classes of specific bacterial poisons. One—typified by diphtheria and tetanus poisons—is produced during the period of energetic growth by the living bacteria, is given off to the surrounding culture fluid as a secretion or excretion, and can be obtained in bacteria-free filtrates at a time when few, if any, of the micro-organisms have died or disintegrated. These are spoken of as “true toxins” or “exotoxins.”

The other group—typified by the cholera poisons as described by Pfeiffer—is apparently an intracellular, constituent part of the bacterial body—not given off during life and not, therefore, obtained in filtrates of young living cultures. If the cultures are preserved until cell death has taken place and the dead bodies have been extracted by the culture fluid, the filtrate becomes gradually more toxic. The bodies of such bacteria are in themselves powerfully toxic when injected, dead or alive. These poisons for obvious reasons Pfeiffer has named the “*endotoxins*,” since he regarded them as specific and definite substances, present as such in the living bacterial cell.

In addition to the endotoxins the bacterial protein contains substances which attract and lead to the accumulation of leukocytes. In other words, they exert a positive chemotactic influence. This was first observed in 1884 by Leber,¹⁵ who induced the formation of pus by injecting dead staphylococcus cultures, and, later, found that the same effect resulted from the injection of alcoholic extracts of staphylococci. These chemotaxis-inducing substances were later particularly studied by Buchner. Buchner¹⁶ extracted them from many varieties of bacteria, independent of pathogenicity. Although there are quantitative differences, all bacteria seem to contain such substances, and Buchner believed the chemotactic property to be a general attribute of the bacterial protoplasm. He speaks of his extracts as *bacterial proteins*.

The true *toxins* or *exotoxins*, then, appear to be products of living bacteria given off from these very much as are the ferments and enzymes by which micro-organisms cause cleavage of carbohydrates or proteins—and indeed the French school, from the first, compared

¹⁴ Pfeiffer. *Zeitschr f. Hyg.*, Vol. II, 1892.

¹⁵ Leber. “Über die Entzündung,” Leipzig, 1884.

¹⁶ Buchner. *Berl. klin. Woch.*, 1890.

these toxins to enzymes, with which, as we shall see, they have much in common. The *endotoxins*—on the other hand—at least as conceived by Pfeiffer, are structural ingredients of the bacterial protoplasm which are toxic when brought into solution as the cells break up.

Concerning the accuracy of this conception, however, much doubt has recently arisen, as a result of researches which will be discussed below.

These two types of poison, moreover, differ from each other not only in mode of origin but in biological characteristics far more fundamental than this.

The discovery of diphtheria toxin by Roux and Yersin was followed by diligent investigations into the toxic properties of all known pathogenic bacteria, and it was soon found that a few only of these germs could produce poisons biologically similar to that found in diphtheria cultures. It was in the course of investigations of this kind, indeed, that Pfeiffer, failing to discover an exotoxin in cultures of cholera and other germs, formulated his endotoxin theory.

The list of true toxin or exotoxin producers, then, is short. Among the more important are, in addition to the diphtheria and tetanus bacilli—which have been mentioned above—the *Bacillus botulinus*,¹⁷ the *Bacillus pyocyaneus*,¹⁸ and that of symptomatic anthrax.¹⁹ It has also been claimed that similar toxins are formed by the cholera spirillum (Braun and Denier),²⁰ by the dysentery bacillus of the Shiga-Kruse type (Kraus and Doerr)²¹ and the *Bacillus typhosus* (Arima).²² In the cases of the three last-named organisms, however, the secretion of a true exotoxin has not been accepted as a fact by all observers. Indeed, even though such substances may possibly be produced by these bacteria in small amounts it is not likely, in the light of our present knowledge, that they play more than a secondary rôle in the toxemic manifestations of cholera, dysentery, and typhoid, the important poisons in these cases being those derived from the bacterial cell bodies.

Similar in essential properties to the true exotoxins also are the erythrocyte poisons (hemotoxins) produced by many bacteria which cause hemolysis of red cells, and the leukocyte-destroying poison (leukocidin) which is a product of the *Staphylococcus aureus*.

All of these "true bacterial toxins" or exotoxins, apart from similarity of origin, as soluble secretions of the living bacteria, possess certain common biological characteristics which sharply differentiate

¹⁷ Kempner. *Zeitschr. f. Hyg.*, Vol. 26, 1897.

¹⁸ Wassermann. *Zeitschr. f. Hyg.*, Vol. 22, 1896.

¹⁹ Grassberger and Schattenfroh. *Wien Deuticke*, 1904.

²⁰ Braun and Denier. *Ann. de l'Inst. Past.*, Vol. 20, 1906.

²¹ Kraus and Doerr. *Wien kl. Woch.*, 42, 1905.

²² Arima. *Centralbl. f. Bakt.*, I, Vol. 63, 1912.

them from the "endotoxins." These characteristics they share with a number of non-bacterial substances such as the vegetable poisons ricin, croton, and abrin, with animal poisons like snake venom and spider poison (arachnolysin), and, in certain important respects, with the substances spoken of as enzymes.

Thus the bacterial true toxins are not biologically unique substances. Both in themselves and in regard to the reactions they elicit when injected into the animal body, they share certain cardinal properties with analogous substances derived from the higher plants and from animals. And it is important to recognize at once that we are dealing here, as in other phases of the study of bacterial immunity, with broad biological laws, which find application not only in bacteriology, but in general pathology and in the phenomena of protein metabolism in general. It so happens that these phenomena have been studied and are most easily elucidated in connection with bacteria. But their general significance must not be lost sight of.

The cardinal characteristic which unites all of these substances into a single well-defined biological group is their property of inducing the formation of *antitoxins* when injected into animals. This property is so important and its thorough comprehension so essential that we may be permitted to digress briefly in order to make it clear.

As we shall see, in subsequent chapters, all substances which lead to the formation of specifically reacting antibodies in the treated animal are spoken of as "*antigens*" or "antibody-inducing substances." The class of "antigens" is a large one, including all known proteins, and possibly some of the higher proteid split products, and protein-lipoid combinations, though the "antigenic" properties of the last two are still in controversy. But among this large group of substances it is only the bacterial true toxins (exotoxins), obtained in broth filtrates of living cultures, together with the vegetable poisons and other substances we have classified with them above, which induce in the blood of the treated animal a neutralizing antibody—(antitoxin)—which inhibits quantity for quantity the activity of the injected toxin or vegetable or animal poison. This property of eliciting the production of antitoxin in the animal body alone separates these substances sharply from all other antigens, toxic or otherwise, and, in this respect, they differ sharply from the so-called "endotoxins" against which no antitoxins can be produced.

As an important secondary characteristic of this group of substances we may regard their chemically indefinable nature. In the case of none of them have we any definite knowledge of chemical constitution except in so far as it has been hitherto impossible to separate them from the protein molecule. The intensive chemical study of the toxins has universally resulted in failure to obtain a protein-free product which has the characteristic toxic properties of

the original filtrate, or its antitoxin-inducing power. Concerning the methods which have been employed in the study of the chemistry of these substances we will have more to say in another place.²³ It is safe to summarize all this work for our present purposes, by stating that, whatever the method employed, until now all of the preparations obtained have given one or another of the protein type-reactions, and that none of them can be positively accepted as protein-free. The results here obtained have been entirely analogous to those obtained in similar investigations upon enzymes. (See also discussion of antigens, chapter 4.)

The analogy with enzymes is indeed a striking one and noted by the first investigators of a true toxin, Roux and Yersin. Biologically, of course, we have the cardinal similarity in that the injection of toxins into animals induces the production of antitoxin, and treatment with enzymes induces specific and neutralizing anti-enzymes. In addition to this, they are alike in their susceptibility to heat (both being destroyed when in solution by temperatures over 80° C.), in their gradual deterioration on standing, and their mysterious activity in small quantities upon disproportionately larger masses of the substances they attack. There is, however, one important difference between the two in their mode of action. For, while the toxins are apparently bound or neutralized by the tissues they attack, the action of an enzyme seems rather to be a process in which the enzyme unites with the substance it acts upon, is released as the result is attained, and freed for further action, without noticeable loss of quantity. Such catalytic properties have not yet been satisfactorily demonstrated for the bacterial toxins. However, there are other modifying factors which may account for lack of similarity in this respect, and in all other important points the two classes of substances are closely analogous.

The property of heat sensitiveness, which is a characteristic of bacterial exotoxins and enzymes, is shared with them by all of the substances mentioned above except snake venoms. Snake venoms are not destroyed completely until the temperature is raised to 75°-80° C. The earlier contention of Leclainche and Vallée, that the toxin of symptomatic anthrax possessed similar heat stability has been satisfactorily refuted by Grassberger and Schattenfroh,²⁴ who find that heating it to 50° C. for an hour completely destroys it.

There is another important attribute of the true toxin which deserves discussion, though we are by no means in a position to offer any satisfactory explanation for it. We refer to the incubation time which elapses between the administration of a toxin and the occur-

²³ An extensive and authoritative summary of this phase of the subject is that of E. Pick in "Kolle u. Wassermann Handbuch," etc., 2d ed., Vol. 1.

²⁴ Grassberger and Schattenfroh. "Über das Rauschbrandgift, etc.," Wien. Deuticke, 1904.

rence of symptoms. Here again snake poisons form an exception—since local manifestations may appear within an extremely short period after the injection of the venom or as the result of a snake bite. However, in the case of all other toxins there is a definite lapse of time between the entrance of the poison and the first symptoms, local or general. This interval is longer when small doses are given—shorter when the doses are large—but is never entirely eliminated—even when many times the fatal dose is given.

In the case of tetanus poison, for instance, injections into a horse may not cause symptoms for as long as four or five days. In mice, animals that are extremely susceptible, the incubation time may be shortened from 36 to 12 hours if we inject 3,600 lethal doses, but, in any case, whatever the dose, this interval cannot be shortened below 8 or 9 hours.²⁵ Many attempts have been made to explain this. Ehrlich, as we shall see, assumes that the action of a poison depends upon two occurrences: one, the union of the poison with the vulnerable cell, the other the gradual injury of the cell by the toxic atom groups in the poison molecule. The time necessary for the institution of this process, he believes, explains the interval. Richet has suggested that the toxin itself may not be potent until acted upon by the body of the recipient and transformed into a potent form. His views are more directly related to the phenomenon of anaphylaxis and are discussed in another section. De Waele has recently advanced a theory which implies that the incubation time represents the period necessary for the gradual concentration of the poisons in the vulnerable tissues, a process which depends either upon chemical affinities or solubility of the toxins in the cell lipoids. A little at a time would then be absorbed by the vulnerable cells as they come in contact with the poison, through the circulation, and the symptoms would not appear until a definite intracellular concentration had been attained. His views are so closely bound up with the theories on the selective action of the toxins upon individual tissues and organs that they will be rendered clear as we proceed with a discussion of the latter.

The majority of pathogenic bacteria do not, as we have seen, produce *true toxins* or *exotoxins*. Cultures of cholera spirilla, plague bacilli, and of many other bacteria do not yield toxic filtrates until the cultures have been allowed to stand for prolonged periods during which extraction and possibly autolysis have occurred. In these cases, moreover, definite toxic properties can be demonstrated in the dead cell bodies or in extracts prepared by various methods. In no case, however, is the injection of these “endotoxins” followed by the production of antitoxins. It was very natural to suppose that in micro-organisms of this class the toxic principle might be present in the form of a preformed intracellular poison which could be ex-

²⁵ De Waele. *Zeitschr. f. Imm.*, Vol. 4, 1910.

tracted or which became free as cell-death occurred and disintegration ensued.

It was assumed that, when bacteria entered the animal body and were destroyed by the action of the serum or cells, these endotoxins were liberated and poisoning resulted. The very protective action of the serum, which prevented the extension of the infectious invasion, by limiting bacterial growth, was thus looked upon as the agency by which the endotoxins were set free. Experiments by Radziewsky and others, in which it was shown that large doses of bacteria injected into immunized animals were violently toxic and more rapidly fatal than corresponding amounts injected into normal animals, were taken to mean that in the immune animals a more powerfully cell-destroying property of the serum led to a more rapid liberation of the endotoxins.

This was the conception of Pfeiffer and, in more recent theoretical discussions, that of Wolff-Eisner. Its essential features consisted in the assumption that the poisons were preformed and were contained within the cell body as such, and that they were specific for each micro-organism, determining to a certain extent its pathogenic properties. Thus typhoid endotoxin, cholera endotoxin, or dysentery endotoxin was supposed each to possess its own particular pharmacological properties by which the clinical manifestations of the respective diseases were partially determined.

It is chiefly the work of Vaughan²⁸ which has begun to throw doubt upon Pfeiffer's original views, in that Vaughan has shown that all proteins, bacterial or otherwise, would yield, upon cleavage with alkalinized alcohol, toxic split products which possessed many of the pharmacological properties of the so-called endotoxins. In fact, Vaughan succeeded in producing, in animals, fever and other symptoms which are generally associated with infection, merely by injecting into them graded quantities of his toxic split products.

Following Vaughan, Friedberger succeeded in showing that toxic substances similar to Vaughan's split products are formed when bacteria of various species are subjected to the action of normal or immune sera, and that such poisons were pharmacologically alike and produced with equal ease from pathogenic and non-pathogenic micro-organisms. These phenomena are discussed in greater detail in our section on bacterial anaphylaxis. It is necessary, however, to point out in this place the uncertainty in which these researches have left the conception of endotoxins. They suggest that the toxic effects following upon the introduction of pathogenic bacteria into the animal body are not due to endotoxins, but are rather the result of the action of toxic cleavage products formed in the reaction between blood plasma and bacterial cell. These split products

²⁸ For a complete discussion of Vaughan's work see Vaughan, "Protein Split Products," Lea & Febiger, Phila. and N. Y., 1913.

are not conceived as specific for individual bacteria but may be formed from all bacterial proteins, both the pathogenic and the non-pathogenic. The differences in pathogenicity between bacteria of this class would then depend entirely upon their powers to invade—not at all upon their possession of individually peculiar cell poisons. The differences in clinical course and toxemic manifestations would be taken to depend entirely upon the accumulation and the distribution of the invading germs, and the consequently variable energy in the production of the toxic split products from them. Considerable experimental evidence has accumulated in favor of this point of view. We will reserve a consideration of this for a later chapter.

In order to do injury to the infected individual the bacterial poisons must be produced in such locations that they can easily enter the physiological interior of the body. None of the poisons that have been so far investigated can produce injury when introduced into the alimentary canal. In this location they are, as a rule, destroyed, or they pass through without doing harm. Neither diphtheria toxin nor tetanus toxin will produce symptoms when introduced intrainstestinally.^{27 28 29 30} Even cholera poison does not pass through the uninjured intestinal wall. Kruse³¹ assumes, and Kolle and Schürmann³² seem to agree with him, that the absorption of cholera poison does not occur until the intestinal wall has been injured by the actual growth of the living bacteria. Kruse calls attention to experiments by Bürgers in which enormous quantities of cholera poison, i. e., 200 cultures of dead or living cholera bacilli, could be administered to healthy guinea pigs and rabbits by mouth without harm in spite of the fact that these animals are definitely susceptible to the poisons and although the poisons are not injured by the intestinal ferments. It is likely therefore that the absorption of poison begins only after the bacteria have extensively invaded the intestinal mucosa and, by injuring tissue, have opened paths for absorption. In the case of diphtheria probably a similar condition exists in that the localized injury to the mucous membrane at the point of lodgment of the primary infection prepares a portal of entry. The poison of the *Bacillus botulinus* alone seems to form an exception to this rule,³³ since this substance, though apparently a true bacterial toxin, is absorbed directly from the intestinal canal. With most bacteria this problem does not arise, since the poisons are

²⁷ Meyer and Gottlieb. "Exp. Pharmacol.," Urban & Schwartzberg, Berlin, 1911.

²⁸ Ransom. *Deutsche med. Woch.*, No. 8, 1898.

²⁹ Nencki. *Centralbl. f. Bakt.*, Vol. 23, 1898.

³⁰ Carrière. *Ann. de l'Inst. Past.*, Vol. 13, 1899.

³¹ Kruse. "Allgemeine Mikrobiologie," Vogel, Leipzig, 1910, p. 934.

³² Kolle and Schürmann in "Kolle u. Wassermann Handbuch," 2d Ed., Vol. 4.

³³ Madsen in "Kraus u. Levaditi, etc.," Vol. 1.

elaborated within the tissues, where resorption is a necessary result.

Like alkaloids and other organic as well as inorganic drugs, the action of many bacterial poisons is largely selective. Most of these poisons may excite inflammatory reactions if concentrated in any part of the body, but, in addition to this, there is a specific distribution after introduction which indicates that the poison goes into selective relationship with certain tissues and cells. This fact is most clearly illustrated by the bacterial hemotoxins which specifically injure the red blood cells of the infected individual and by such substances as the leukocidin produced by the *Staphylococcus aureus*, a poison which directly and visibly injures the white blood cells. Here the action is specifically aimed at a well-defined variety of body cell.

In considering this problem in connection with infectious disease, it is of great importance to distinguish between selective injury by the poisons transported through the body by the lymph, blood, and other channels, on the one hand, and the selective lodgment of the micro-organisms themselves on the other. The latter may occasionally depend on local cultural advantages for the particular bacteria in one organ or another, but may just as often be determined by the peculiar manner of entrance to the body which is most suitable for lodgment of the germs in question, and the degree of local resistance at the point of entrance, which determines whether or not the infection shall be locally limited or permitted to invade beyond this point. In the case of a disease like acute anterior poliomyelitis, where our knowledge of the micro-organisms which cause the disease is yet in its infancy, it is impossible to decide whether the injuries noted in the motor areas of the cord and medulla are due to toxins or the lodgment of the germs themselves. In the case of rabies it seems reasonably sure that the micro-organisms themselves select the nervous system. In such instances as the injury of the motor areas by tetanus poison, that of certain peripheral nerves by diphtheria toxin, or even the characteristic lesions of post-syphilitic maladies like tabes, we can be reasonably sure that we are dealing with the specific action of the poisons, independent of actual localized growth of the infectious agents.

Diphtheria toxin, after distribution through the body, may act upon many different tissues, as is evident by degenerations in the heart muscle, liver, and kidney, and the petechial hemorrhages in serous surfaces. In addition to this general action, however, there is a very marked selection of certain nerve centers. By Meyer and Gottlieb³⁴ diphtheria toxin is classed as a specific vascular poison. Its action results in a rapid sinking of the blood pressure with final

³⁴ Meyer and Gottlieb. "Pharmacology Trans. Halsey," Lippincott, 1914, p. 556.

cardiac death in spite of artificial respiration. These manifestations seem to have a central origin, with particular action upon the vagi and the phrenic nerves. Apparently also the localization of the diphtheritic lesion may influence the selection of individual nerves, the most concentrated action taking place upon the nerves whose endings are distributed in this particular region, for, as Meyer and Ransom³⁵ have shown, this poison, like tetanus toxin, may be absorbed into the nerves directly through the nerve endings. An interesting selective action also of diphtheria poison is the apparently specific alteration of the suprarenal glands which is regularly noticed, as enlargement and congestion, in diphtheria-infected guinea pigs, and which has been associated by many workers with the characteristic drop in blood pressure which accompanies all severe cases of the disease. Abramow³⁶ has studied this lesion particularly, and believes that it consists in a degeneration and final disappearance of the chromaffin substance and of the medullary cells. He believes that this, together with degeneration of the heart muscle itself, is of great importance in causing the characteristic vascular failure.

In botulinus poisoning there is, as Marinesco³⁷ and Kempner and Pollack³⁸ have shown, a direct effect upon the cells of the anterior horns with degenerative changes in the Nissl granules.

Tetanus poison, which has been studied extensively by pharmacologists, shows a very marked affinity for the nervous system, as, in fact, the symptoms of tetanus indicate. Indeed, while many of the bacterial poisons are distributed by the blood stream to the point of final attack, in tetanus the absorption of the toxin from the lesion or the point of injection takes place entirely by the path of the nerves.

That this method of poison distribution might be, among others, an important one was suggested as early as 1892 by Bruschettiini,³⁹ who found tetanus toxin in the nerves but not in the adjacent muscle and other tissues surrounding the point of subcutaneous injection. Similar results were obtained subsequently by Hans Meyer, whose experiments were confirmed and extended by Marie and Morax.⁴⁰ Finally Meyer and Ransom⁴¹ furnished complete proof that the poison was absorbed from the blood and tissues by the peripheral nerve endings alone and was transported centripetally only by the paths of the neurons. The experimental facts elicited may be summarized as follows:

³⁵ Meyer and Ransom. *Arch. de pharmacodyn.*, Vol. 15, 1905, also Meyer, *Berl. klin. Woch.*, 25 and 26, 1909, also *Arch. f. exp. Path. u. Ther.*, Vol. 60, 1909.

³⁶ Abramow. *Zeitschr. f. Imm.*, Vol. 15, 1912.

³⁷ Marinesco. *Compt. rend. de la soc. de biol.*, Vol. 3, 1896.

³⁸ Kempner and Pollack. *Deutsche med. Woch.*, 32, 1897.

³⁹ Bruschettiini. *Riforma medica*, 1892.

⁴⁰ Marie and Morax. *Ann. de l'Inst. Past.*, 1902.

⁴¹ Meyer and Ransom. *Archiv f. exp. Path. u. Pharm.*, 49, 1903.

1. When tetanus toxin is injected into the thigh muscles of a guinea pig the poison is found at first only in the sciatic nerve of the same side and in the blood. (The determination of poison was made by injecting macerations of the respective tissues into mice.) If examination was delayed until the symptoms had become generalized, the poison was found in the opposite sciatic, but the muscle bundles, fat, etc., from the vicinity of the injection area were poison-free.⁴²

2. When a nerve is cut poison absorption ceases as soon as axis cylinder degeneration has set in.

3. If the nerve is cut before the poison is injected the distal end contains poison, the proximal end does not. This again shows that the nerve absorbs the toxin not from its capillaries but solely through the end organs.

4. If a nerve which already contains poison is severed, toxin will disappear rapidly from the proximal end, since it no longer obtains a renewed supply from the periphery.

5. If antitoxin is injected into the nerve, above the point of injection, it will successfully bar the way for the ascending toxin.

6. Severing of the spinal cord prevents the passage of the poison from below upward.

These facts ascertained in the case of tetanus find their parallel in the phenomena of the distribution of rabic virus,⁴³ as well as in that of poliomyelitis, in both of which there seems to be a progressive centripetal transportation through the nerves. However, in these conditions we are probably dealing not with a poison but with a living virus and, though analogous, the conditions are not entirely comparable.

From the practical point of view these facts regarding tetanus may explain the frequent failure of therapeutic success attending the injection of tetanus antitoxin after the symptoms of the disease have set in, since in such cases the poison is already distributed to the nerves and is largely inaccessible to the antitoxin. They also have pointed a way toward a more hopeful therapy, namely, the method of injecting the antiserum directly into the nerves about the point of injury. It is not surprising, however, in view of the stated facts, that even this is unsuccessful when done at too late a time, after a considerable amount of poison has already passed above the point of injection to the spinal centers.

Such selective action on the part of the bacterial poisons is entirely analogous to the similar specific action of alkaloids, narcotics,

⁴² In view of our discussion of the importance of fats in the absorption of tetanus toxin, it seems inconsistent that the toxin does not concentrate in fatty as well as in nervous tissues. This Meyer explains by the inactive and poorly vascularized condition of the fat tissues.

⁴³ Di Vestea and Zagari. *Fortschr. d. Med.*, Vol. 6, 1888.

and other drugs. In order that the poison may act upon a cell we must, of course, assume that it has either chemical or physical affinity for this cell. The problem, as many writers have pointed out, is strongly analogous to that of tissue staining. A dye must be able to form a chemical union with the cell or it must be soluble in the cell substance in order to stain it. The chemical difference between cells is a delicate one and not often definable by our present methods. We can obtain an insight into the principles probably underlying selective action only by inference from the relation between the chemical constitution of drugs or their physical properties, solubility, etc., and their respective tissue affinities. These problems are difficult and, to a large extent, obscure. They cannot be directly investigated upon bacterial poisons since these are themselves of chemically unknown nature. But the study of drugs of known constitution has revealed certain definite relations of this kind which have furnished analogies from which the general principles of selection in bacterial poisons can be surmised.

It is a well-known fact to pharmacologists that there is a definite relation between chemical structure and toxicity. Fraenkel⁴⁴ expresses it as follows: "By the addition of identical atom groups in an identical manner, similarly acting substances are obtained." He cites the well-known example of curare; whichever the path by which this poison is injected it leaves intact the tissues with which it comes in contact, but after general distribution acts specifically upon the nerve endings. It had been discovered by Brown and Fraser⁴⁵ that by introducing methyl radicles (CH_3) into molecules of various alkaloids, strychnin, morphin, atropin, and others, substances were obtained which paralyzed nerve endings, and this irrespective of their previous physiological action. It appears that the combination of four methyl radicles attached to the nitrogen atom (quaternary bases) universally possesses this paralyzing action. Tertiary bases on the other hand lack this property.



"Ammonium base"



"Tertiary base"

⁴⁴ Sigmund Fraenkel. "Arzneimittel Synthese," 2d Ed., Springer, Berlin, 1906.

⁴⁵ Brown and Fraser. *Trans. Royal Soc. of Edinburgh*, 25, 1868, cited from Fraenkel.

Quaternary

Subsequently Böhm^{46 47} discovered that curare contains two bases—the one, “curin,” is slightly toxic and is a tertiary base; the other, which possesses the typical curare action, “curarin,” is an “ammonium base.” By “methylizing” curin, curarin could be obtained.

From these and other examples it is clear that in a certain number of cases actual chemical affinity must play a part in toxic action; on the other hand, there are many cases in which toxic action seems to depend merely upon physical conditions such as solubilities. Meyer and Overton's well-known theory of narcosis maintains that certain narcotics exert their action by passing out of blood and lymph solution into solution by the fat-like, lipoidal substances (lecithin, cholestrin, etc.) contained in the nerve cells, because the latter are better solvents for them than is the blood plasma. This theory of Meyer and Overton has stimulated much investigation and speculation, and it is not unlikely that it is valid in the case of many narcotics, although it does not explain the action of narcotics in general; for Dickson notes that chloral hydrate, for instance, is more soluble in water than in oils, and some narcotic drugs like alcohol exert definite action on proteins and are oxidized in the body. These are pharmacological questions of which we cannot speak with authority. We wish merely to point out that the action of poisons upon the body may depend in some cases upon mere physical or mechanical relationship between the two.^{48 49}

As regards bacterial poisons the union between poison and susceptible cell is extremely firm and difficult to dissociate in many instances, and this points to the possibility that, in these cases at least, true chemical union takes place rather than merely a loose combination like that of the solution of one substance in another. Furthermore, the complete inactivation of some poisons by mixture with the cells of tissues capable of binding them would likewise point to more than mere physical union. Nevertheless, it does not by any means exclude the thought that the poisons may, in fact, go into selective relationship with special cells because of physical properties, such as solubility in the lipoidal cell membranes,^{50 51} and may

⁴⁶ Böhm. *Arch. de Pharm.*, cited from Fraenkel.

⁴⁷ See also Dickson, “A Manual of Pharmacology,” E. Arnold, London, 1912.

⁴⁸ Ivar Bang. “Biochemie der Lipoide,” Bergmann, Wiesbaden, 1911.

⁴⁹ Meyer and Gottlieb. “Experimentelle Pharmakologie,” 2d. Ed., Urban & Schwartzberg, Berlin, 1911.

⁵⁰ For Overton's theory of osmosis see R. Höber, “Physikalische Chemie der Zelle u. Gewebe,” Leipzig, Engelmann, 1911.

⁵¹ Compare also, regarding this entire question, the discussion in P. Th. Müller, “Vorlesungen über Immunität, etc.,” Fischer, Jena, 1910.

subsequently be bound chemically or destroyed by oxidation or enzymotic hydrolysis after such entrance. In such a case the actual specificity would yet depend on purely physical properties.

In addition to the specific physical and chemical affinities between the poisons by certain cells there are probably also certain fortuitous factors connected with the distribution and local accumulation of the poisons which have some weight in determining the location of injury. For the specific selection is not absolutely strict and there are probably few parenchyma cells in the body that are entirely insusceptible to injury if the poisons are sufficiently concentrated upon them. Thus, to cite an analogy from the toxicology of non-bacterial poisons, in lead poisoning, as Meyer and Gottlieb point out, the paralysis of the extensors of the arm occurs chiefly in adults who use these muscles in the exercise of their professions (painters, type-setters), while in children and in animals, in which no such selective use of particular muscle groups is habitual, lead paralyses are atypical, attacking legs as well as arms. It is not unlikely that the frequent injury of the heart muscle by bacterial poisons or the irregular parenchymatous changes in various organs is determined by analogous fortuitous factors, in that functional activity and increased metabolism may predispose to injury.

Bacterial poisons also may produce their lesions in the course of excretion. This seems likely in the case of typhoid poisons in which we have often seen bloody diarrhea in rabbits within a few hours after intravenous injection of powerfully toxic culture filtrates. In connection with the dysentery bacillus Flexner and Sweet⁵² have studied the conditions carefully. They succeeded in showing first that the introduction of the dysentery poison into the lumen of the intestine does no harm and that the toxin is slowly destroyed by peptic and tryptic digestion. They concluded that probably no absorption of the poison through the uninjured intestinal mucosa takes place. They then showed that the toxin after intravenous administration is excreted by the intestine and that the inflammatory reactions and injury of the mucosa are incident to this act of elimination.

Whether or not the kidneys are injured in the same way it is difficult to decide. In many infectious diseases, of course, the bacteria themselves pass through the kidney into the urine, and renal injury may result from the actual presence of the bacteria in the kidney; however, renal injury may also occur without this, and it is not at all impossible that the conditions here are similar to those just described for the intestine.

All the facts which we have considered indicate that, although most bacterial poisons can injure many different tissues, yet in some cases there is a particular susceptibility on the part of an individual

⁵² Flexner and Sweet. *Jour. of Exp. Med.*, Vol. 8, 1906.

tissue which is independent of accidental factors and seems to be due to specific chemical or physical affinity. It seems even that in tetanus, botulismus, and a few other conditions there is a differential selection of particular areas within a tissue like the nervous system, just as this occurs in the case of certain drugs. As stated above we have no satisfactory scientific explanation for this, but a great deal of work has been done to show that the bacterial poisons actually unite with and are taken up by the susceptible tissues.

Indirectly, proof of this has been brought by the demonstration of the rapid disappearance of various toxins from the blood streams of susceptible animals and their persistence in the circulation of animals insusceptible to them. Thus Dönitz⁵³ has shown that tetanus toxin injected into the blood stream of a susceptible animal rapidly diminishes in quantity, and Knorr,⁵⁴ in similar experiments, showed that the demonstrable disappearance of such toxins out of the blood stream is synchronous with the appearance of symptoms, a fact which excludes disappearance by excretion. Conversely Asakawa⁵⁵ showed that in pigeons, which are but slightly susceptible, tetanus poison could be demonstrated in blood, liver, spleen, kidneys, and muscles six days after injection, but not in the brain, showing that in this organ, at least, there must have been either a union or a destruction of the poison. Similar to these results are those of Metchnikoff,⁵⁶ who found the poison unchanged after two months in the circulation of insusceptible animals (lizards).

Direct evidence of union between susceptible tissues and poison has been furnished by the experiments of Wassermann and Takaki,⁵⁷ who showed that the brain and cord tissues of rabbits and guinea pigs, mixed with tetanus toxin before injection, served to neutralize its harmful effects. And it appears that the toxin-neutralizing property of the brain substances of various animals is proportionate to their individual susceptibility to the poison. Thus Metchnikoff⁵⁸ not only confirmed the results of Wassermann and Takaki for rabbits and guinea pigs, but showed further that the brains of chickens, animals that are but moderately susceptible, possess a correspondingly slighter neutralizing power, and, further, that brain tissues of entirely insusceptible cold-blooded animals, turtles and frogs, possess absolutely no neutralizing properties.

The original interpretation by Wassermann of these facts was based on the assumption that the poison was bound to the brain tissue

⁵³ Dönitz. *Deutsche med. Woch.*, No. 27, 1897.

⁵⁴ Knorr. *Fortschr. der Medizin*, 1897, No. 17, and *Münch. med. Woch.*, 1898, Nos. 11 and 12.

⁵⁵ Asakawa. *Centralbl. f. Bakt.*, Vol. 24, pp. 166 and 234.

⁵⁶ Metchnikoff. "L'Immunité dans les maladies Infect.," Paris.

⁵⁷ Wassermann and Takaki. *Berl. klin. Woch.*, 1898, No. 1.

⁵⁸ Metchnikoff. *Ann. de l'Inst. Past.*, 1898, p. 81.

just as it is bound to antitoxin. Experiments by Besredka⁵⁹ have cast some doubt upon this. This worker's experiments seem to indicate that a brain emulsion which has been saturated with the toxin can be rendered capable of absorbing more toxin if tetanus antitoxin is mixed with it. In other words, the affinity of the antitoxin for the toxin is stronger than that of the brain substance for the poison, and that the union toxin-brain tissue is very easily dissociated; as indeed it should if the union were purely a physical one depending on solubility.

After it had been shown that the poisons which acted specifically upon certain cells were actually taken up by these cells, a number of attempts were made to determine chemically the tissue element which united with the poisons. Noguchi⁶⁰ showed that cholesterin and alcoholic extracts of blood serum neutralized tetanolysin. The same thing was later shown by Müller,⁶¹ and Landsteiner⁶² showed that ether extracts of red blood cells likewise neutralized this poison. In a later study by Landsteiner and von Eisler⁶³ the relation of the tissue lipoids to various toxic substances was still more definitely established. They studied first the various hemolysins and found that extraction of blood cells with ether rendered the stromata less capable of binding the hemolytic substances. The same thing they showed for bacteriolysins, in the latter case demonstrating at the same time that the ether extracts of bacterial bodies possessed slight binding properties for the bactericidal substances of the serum. These experiments have, of course, a merely indirect significance in the present connection, since they do not deal with the type of poisons we have discussed. However, Landsteiner and von Eisler also worked with tetanus toxin and found that the treatment of the brain substance of guinea pigs with ether, by taking out lipoidal substances, considerably reduces the power of this tissue to bind and neutralize the tetanus poisons.

Takaki,⁶⁴ who investigated these relations in great detail, isolated an alcohol-soluble element, cerebrin, from nerve tissues, a substance to which he ascribes the toxin-binding properties. Overton and Bang⁶⁵ found, furthermore, that cholesterin and lecithin inhibit the action of cobra venom, a poison which is in so many ways similar to those produced by bacteria. Taking into consideration all available evidence, we are forced to admit that the lipoids seem to play an important rôle in determining the selective action of the nervous sys-

⁵⁹ Besredka. *Ann. Past.*, 1903, p. 138.

⁶⁰ Noguchi. *Univ. Pa. Med. Bull.*, Nov., 1902.

⁶¹ Müller. *Centralbl. f. Bakt.*, Vol. 34, 1903.

⁶² Landsteiner. *Wien. kl. Rundschau*, 13, 1905.

⁶³ Landsteiner and von Eisler. *Centralbl. f. Bakt.*, 39, p. 318, 1905.

⁶⁴ Takaki. *Beitr. zur chem. Phys. u. Path.*, 11, No. 19, 1908.

⁶⁵ See Ivar Bang, "Biochemie der Lipide," Bergmann, Wiesbaden, 1911.

tem by the bacterial poisons. It may not, of course, be an influence depending merely upon the solubility of the harmful substances in the lipoids themselves. For, as Bang expresses it, "the lipoids possess to a high degree the property of altering by their presence the solubilities of other bodies," and it is quite possible that in the tissues they are present as lipoid-protein combinations. Their action in determining the solubility of toxins in a given cell may therefore be a purely indirect one.

It is of some interest in this connection to recall the experiments of De Waele,⁶⁶ which bring out another clear analogy between alkaloids and bacterial poisons in their relation to lecithin. He found that the addition of small quantities of lecithin increases the activity of both toxins and alkaloids in the animal body, whereas larger amounts inhibit both.

⁶⁶ De Waele. *Zeitschr. f. Immunit.*, Vol. 3, 1909, p. 504.

CHAPTER III

OUR KNOWLEDGE CONCERNING NATURAL IMMUNITY, ACQUIRED IMMUNITY, AND ARTIFICIAL IMMUNIZATION

NATURAL RESISTANCE AGAINST INFECTION

IN the preceding chapters we have confined ourselves largely to the consideration of those properties of the bacteria which determine their ability to infect. In this discussion, however, we have repeatedly emphasized the fact that every infectious disease is the result of a struggle between two variable factors—the pathogenic powers of the bacteria on the one hand, and the resistance of the subject on the other, each of these again modified by variations in the conditions under which the struggle takes place. Thus a given micro-organism may be capable of causing fatal infection in one individual but may be only moderately virulent or even entirely innocuous for another. Conversely the same individual may be highly susceptible to one variety of bacteria, but highly resistant to others. Even in reactions with one and the same micro-organism, the susceptibility or resistance of the individual may be determined by variations in the physiological state or by the environmental conditions under which the two factors—invader and invaded—are brought together. Therefore, the conceptions “resistance,” “immunity,” and its opposite “susceptibility,” are relative terms which can never be properly discussed without careful consideration of all modifying conditions which influence them.

The science of immunity deals with a detailed analysis of these variables. Its ultimate practical aim is the determination of methods by which an original susceptibility can be transformed into resistance or even immunity. And the rational method of approaching this subject consists in a careful study of the conditions of susceptibility and immunity as they exist naturally in the animal kingdom.

The mere fact that both animals and man are in constant contact with infectious micro-organisms, many of them in a high state of virulence, indicates in itself that the animal disposes normally over a defensive mechanism of considerable efficiency.

To a certain extent, of course, this escape from harm is due to

the external defences of skin and mucous membrane which, in the healthy state, mechanically prevent the entrance of the micro-organisms into the body. For we have seen, in another place, that few of the bacteria can pass through the uninjured surfaces. Moreover, added to this, there is some protection in the bactericidal properties of the secretions. An example of this is the inhibitory power exercised by the acidity of the normal gastric juice upon the cholera spirillum. In order to infect the intestinal canal of guinea pigs with these organisms Koch found it necessary to neutralize the gastric juice with sodium carbonate solutions, and other observers have found it necessary to inject directly into the duodenum. But even after entrance into the animal tissues a second line of defence is normally encountered by all invading germs which tend to inhibit their further progress more or less perfectly. This active opposition to the bacteria after their entrance is expressed chiefly in the anti-bacterial (bactericidal) activity of the blood serum, and the phagocytic powers of leukocytes and other cells. To a certain extent these forces are active against all bacteria in all animals, but they may vary in different species, races, or even individuals in potency against any given infectious agent, and, to a certain extent, variations in resistance may be referable to this. The analysis of these forces, both in the normal and in the artificially immunized animal, forms the substance of the systematic discussions which are to follow, and, for the present, we will confine ourselves to an examination of the facts that have been gathered regarding the actual differences in normal resistance or "Natural Immunity" between various species of animals.

And if we glance over the list of diseases to which different species and races of animals are victim, it is immediately evident that some animals are never spontaneously infected with many of the micro-organisms that cause extensive and fatal ravages in others. Also, within the same race or species, an epidemic sweeping through a community will kill many individuals and leave others unscathed. Such differences point to variations in the defensive mechanism, since the invader in these cases is the same. We speak, therefore, of Natural Immunity which is an attribute of species, that which, within the same species, is racial, and that which, within the same race, is individual. And the attempts to discover the causes underlying such differences in natural resistance have elucidated many of the fundamental principles of immunity in general.

Instances of natural immunity which appear to depend on species are common. We have pointed out, above, that in order to make infection at all possible, it is necessary that the invading germ shall find suitable cultural conditions in the body of the host. It is this simple principle which probably explains the fact that bacteria which cause disease in warm-blooded animals cannot, as a rule, cause dis-

ease in those that are cold-blooded, and *vice versa*. Thus frequent attempts to produce anthrax in turtles, frogs, and other cold-blooded species have failed. Also among warm-blooded animals differences in body temperature have been shown to influence susceptibility. Thus avian tuberculosis does not develop in mammals, nor do the human and bovine types of tubercle bacilli infect birds. And this is probably due to the fact that the avian bacillus has become adapted to growth at from 40° to 45° C., about the normal temperature of birds, while the mammalian bacilli cease to grow when the temperature is raised above 40° C. Another observation which clearly illustrates the influence of body temperature upon susceptibility is that made by Gibier¹ upon anthrax. Frogs are ordinarily resistant to this disease. When they are kept in water at 35° C. a fatal infection can be produced. Nuttall's² experiments with plague infection in lizards illustrate the same point. Kept at 16° C., no infection could take place. Warmed to 26° C., they could be readily infected. It is ordinarily assumed that these results are explicable upon the basis of purely cultural and temperature considerations. And this, indeed, is most likely. It is possible, however, that an additional factor involved in this may be the lowering of the general resistance of cold-blooded animals when warmed, just as warm-blooded animals can be rendered susceptible by chilling.

It is for similar simple cultural reasons, possibly, that diseases which occur spontaneously in carnivora do not occur in purely herbivorous animals. The relative resistance of dogs to anthrax and to tuberculosis may possibly be accounted for in this way. However, there are many micro-organisms which infect easily both carnivorous and herbivorous animals, and it may well be that the frequently cited cases we have mentioned above depend on factors more complicated than mere cultural conditions incident to metabolic differences. In most cases of species resistance, indeed, simple nutritional conditions alone do not serve as valid explanations.

Species resistance may be so perfect that it amounts to an absolute immunity. This is apparently so in the cases cited above, namely the immunity of the cold-blooded species to certain diseases of warm-blooded animals. However, such examples are exceptional. When we are dealing with diseases of warm-blooded animals only, natural resistance, in all but a limited number of cases, is sufficient only to prevent the spontaneous occurrence of the particular disease, or to prevent infection when experimental inoculation with moderate doses is practiced upon normal animals. In most of these cases, however, when the dose experimentally administered is excessive, or the resistance is lowered artificially, by chilling or by any other

¹ Gibier. *Compt. rend. de l'acad. des sc.*, Vol. 94, 1882.

² Nuttall. *Centralbl. f. Bakt.*, Vol. 22, 1897.

form of local or general injury, infection can be accomplished. In the case of protozoan diseases species adaptation is much more rigid and parasites that infect one species are very often restricted entirely to that class, being unable to infect any other animal, even though no striking difference in temperature or metabolism exists.

We may convey the clearest conception of all such species differences by a tabulation of some of the more important infectious diseases of man with a statement in each case concerning its transmissibility to animals, as follows:

Tuberculosis, human type, spontaneously infects man. It is very often observed in monkeys kept in captivity. Cattle, swine, and sheep are probably never spontaneously infected; guinea pigs are highly susceptible to experimental inoculation. Cattle, swine, sheep, and rabbits are relatively very resistant to experimental infection. Dogs and goats are still more so. Birds seem to be entirely refractory.

Tuberculosis, Bovine Type.—Spontaneous infection occurs in domestic animals, chiefly cattle; it is less frequent in sheep, hogs, and horses; it has been reported in dogs and goats. In man infection does occur, but only a small percentage of human tuberculosis is of the bovine type, and these cases are almost exclusively in children. In tabulating 1,042 cases which have been carefully studied, Park and Krumwiede³ report the following figures:

Cases of Tuberculosis in Man (1042)

Over 16 years

Human type 677, bovine type 9.

5 years to 16 years

Human type 99, bovine type 33.

Under 5 years

Human type 161, bovine type 59.

The large majority of bovine infections were abdominal or involved cervical lymph nodes.

Experimental infection is successful in rabbits and guinea pigs, both of these animals succumbing more rapidly to this than to the human bacillus. In fact, the relative resistance of rabbits to the human bacillus is such that rabbit inoculation is one of the most important methods of differentiating between the two types. Birds are refractory.

Tuberculosis of the avian type occurs spontaneously in birds. It may be experimentally produced in rabbits (Strauss and Gamaleia). Injected into cattle it causes a local reaction only.

Tuberculosis of cold-blooded animals is not transferable to warm-blooded animals.

Syphilis spontaneously occurs in man only. It can be inoculated

³ Park and Krumwiede. *Jour. of Med. Res.*, Vol. 23, 1910.

into chimpanzees, in which primary and secondary lesions develop, corresponding mildly to human syphilis. Primary lesions can be produced in lower monkeys. It can be transferred by intratesticular inoculations to rabbits.

Gonococcus infection occurs spontaneously in man only. No typical lesions can be produced in experimentally inoculated animals, though death can be caused by large doses, probably by toxic action.

Influenza bacillus spontaneously infects man only. Experimental infection is partly successful in monkeys only. (Pfeiffer and Beck, *Deut. med. Woch.*, 1893.)

Glanders.—Spontaneous infection occurs in horses and mules; less frequently in sheep, goats, and camels. This disease, like plague, may be regarded as primarily a disease of animals, but man may be infected by direct or indirect contact with the diseased animal. All domestic animals may be infected experimentally with ease, except cattle and rats, in which cases large doses are necessary. Birds show local reactions only. (Wladimiroff—in “Kolle und Wassermann Handbuch,” Vol. 5, 2d Ed.)

Plague occurs spontaneously chiefly in man and in rats. It has also been found in California ground squirrels and in hogs during plague epidemics in Hong Kong. It is highly infectious for guinea pigs and white rats—slightly less so for mice; rabbits are much less susceptible than guinea pigs. Dogs, cats, and cattle are relatively resistant. Birds appear to be immune. Cold-blooded animals are immune unless artificially warmed. (See above.)

Malta fever occurs spontaneously in man and in goats. It is pathogenic for all mammals, but it is not fatal for lower animals when the organisms are directly cultivated out of the human body.

Diphtheria occurs spontaneously in man only. Experimental inoculation is fatal in guinea pigs, rabbits, dogs, cats, and birds. Rats and mice are highly resistant. The typical pseudomembranous inflammation can be produced in susceptible animals only after previous injury of the mucous membrane, and then it rarely shows any tendency to spread.

Tetanus is spontaneous in man, horses, cattle, and sheep. It is found rarely in dogs and goats. Birds are highly resistant to experimental inoculation.

Anthrax is primarily a spontaneous infection of cattle, sheep, and horses; it occurs in man largely through direct or indirect contact with these animals. Guinea pigs, rabbits, and white mice are very susceptible to experimental inoculation. Rats and hogs are less susceptible, and dogs are relatively resistant, though they can be regularly killed by moderate doses intravenously injected. Birds and cold-blooded animals are highly resistant.

Asiatic cholera develops spontaneously in man only. Rabbits

and guinea pigs can be killed by injections of cultures, but die probably of toxemia. In rabbits a cholera-like condition has been produced by injection of the spirilla into the duodenum after ligation of the common bile duct. (Nikati and Rietsch, Ref. in *Deut. med. Woch.*, Vol. II, 1884, p. 613.) Ordinarily no multiplication takes place in the animal body. Pigeons are insusceptible, a fact which helps to distinguish this organism from *Spirillum metchnikovi* and other similar bird-pathogenic spirilla.

Typhoid fever occurs spontaneously in man only. It has recently been produced in a mild form in chimpanzees. Animals are susceptible to the endotoxins and can therefore be killed by injections of bacilli and extracts, but the organism is not invasive as in the case of the lower animals. Typhoid septicemia can be produced in rabbits by inoculating them with especially virulent cultures of the bacilli, or cultures previously grown on rabbit-blood agar (Gay). The typhoid-carrier state may ensue for considerable periods in such animals.

Pneumococcus infection in various forms occurs spontaneously in man. Rabbits, mice, and guinea pigs are highly susceptible. Rats, dogs, cats, cattle, and sheep are relatively resistant.

Staphylococcus and streptococcus infections may occur in almost all of the warm-blooded animals, chiefly as abscess producers. In horses a severe form of pleuropneumonia is caused by them.

Leprosy occurs spontaneously in man only. Lesions simulating human leprosy have been produced in monkeys by inoculation, and partially successful experiments have been made upon the Japanese dancing mouse. Other animals are immune.

Scarlet fever occurs spontaneously in man only. Monkeys may possibly be susceptible, though not all observers have been successful in such experiments. (Draper and Handford, *Journ. of Exp. Med.*, Vol. 17, 1913.) Landsteiner and Levaditi (*Ann. Past.*, Vol. 25, 1911) have succeeded in producing the disease in the chimpanzee, though they failed with lower monkeys.

Small-pox occurs spontaneously in man only. It is probably identical with cow-pox. (See reasons for this assumption given by Haeussler as cited by Paul in "Kraus and Levaditi Handbuch," etc., Vol. 1.) It can be experimentally produced in monkeys.

Measles develops spontaneously only in man. *Macacus rhesus* has been successfully inoculated by Anderson and Goldberger (U. S. Pub. Health Reports, 26, 1911). Other animals are immune.

Typhus fever occurs in man only. Experimentally it has been produced in chimpanzees, *Macacus*, *Cercopithecus*, *Ateles*, and *Myrcetes* monkeys. Anderson has succeeded in producing temperature reactions in guinea pigs by injecting blood from typhus patients or from other similarly infected guinea pigs. More exact information concerning this disease will probably be available soon, if the re-

ported cultivation of the organism of the disease by Plotz is authenticated.

Yellow fever up to the present has been observed in man only.

Poliomyelitis is spontaneous in man only. Can be transmitted to monkeys and—in a doubtful form—to rabbits. No other animals are known to be susceptible.

The above represents an incomplete tabulation of the variations in susceptibility in the animal kingdom for infections which occur spontaneously in man. They will illustrate sufficiently, however, the facts of variable species susceptibility as we have stated them. We might, with equal profit, tabulate the infections occurring spontaneously in any single species of animal and show how variable would be their pathogenic powers for other animals and for man. Thus man is immune to the organism which causes cattle plague, and to that of chicken cholera, and probably to many other diseases peculiar to animals, though, of course, in the case of infections of the human being we are entirely dependent for such information upon observed immunity to spontaneous infection, and upon a few instances of accidental inoculation.

In regard, also, to differences of susceptibility between various *races*, within the same species, many interesting facts have been observed. Thus gray mice are, as a rule, more resistant to streptococcus and pneumococcus infection than are white mice. Algerian sheep are said to be more resistant to anthrax than are European sheep. Of black rats inoculated by Müller⁴ with anthrax over 79 per cent. survived, while of white rats similarly inoculated only 14 per cent. survived.

In man, too, racial differences are marked. The extraordinary susceptibility of the negro to tuberculosis is familiar to all American physicians, and it is well known that Eskimos transported to temperate climates and civilized conditions are particularly prone to contract this disease. Small-pox is considered a relatively mild disease in Mexico. Dr. James Carroll⁵ stated that whites are more susceptible to yellow fever than are negroes, and that among the latter those living nearest the equator are less susceptible than are the more northern races. There seems to be no doubt about the actual occurrence of such racial differences, although, as Hahn⁶ very justly points out, many instances formerly regarded as racial differences of susceptibility may have been simulated by racial, or often religious, differences of custom that influence sanitary conditions, and consequently the incidence of epidemic disease.

Apart from the explanations furnished in a few instances by

⁴ Müller. *Fortschr. der Med.*, 1893. Cited from Sobernheim, in "Kolle u. Wassermann Handbuch," 2d Ed., Vol. 3.

⁵ Carroll in "Mense, Tropenkrankheiten," Vol. 2, p. 124.

⁶ Hahn in "Kolle und Wassermann's Handbuch," Vol. 1.

gross physiological differences such as body temperature, the factors determining species resistance are largely a mystery, and in the matter of racial variations, of course, we have no instances in which such very obvious physiological factors play a part. In attempting to find causes for differences of resistance or susceptibility in general, the nature of the problem makes it necessary for us to examine it from a number of different points of view. A micro-organism may be infectious for a given species of animal more than for another, because of special adaptation to the conditions, nutritive and otherwise, encountered in the tissues of these animals. Such adaptation is illustrated in the experience of Pasteur with "rouget" and with rabies, where passage through one variety of animal enhanced the virulence for this species but reduced it for others; and the same thing is easily demonstrated in the laboratory with so many bacteria that it may be accepted as a principle underlying enhancements of virulence in general. This adaptation implies that, to a certain extent, the part played by the animal body in determining its own susceptibility is passive. *Gonococcus*, for instance, infectious for man only, requires human protein for growth, at least in its first generations outside the body. Its ability to cause disease in man may be largely dependent upon its cultural need of human protein. The resistance of other animals to this disease, then, is, in part, due to their failure to supply proper nutriment. This, as Kolle points out, is analogous to *Atrepsie*, a term used by Ehrlich, in speaking of the insusceptibility of one species to cancerous growths originating in another.

Again, "adaptation" on the part of the bacteria may imply, not only an increased ability to meet altered cultural conditions, but an actual acquisition of greater offensive or invasive powers with which to meet the particular defences opposed to it by the given animal. Thus the increased virulence of typhoid bacilli after cultivation in immune sera would point toward an increased ability to survive under the adverse conditions encountered in the animal body. An organism may possibly acquire particular infectiousness for one species to the exclusion of others, by a succession of spontaneous inoculations—comparable to the experimental passage of the micro-organism through animals of the same species. This is especially probable in diseases such as gonorrhea, syphilis, and some others where infection is usually direct from one person to another. And it is these diseases particularly in which infectiousness is rather strictly limited to the human species.

Regarding the matter purely from the point of view of the animal body and the factors which determine its powers to ward off a given infection, we may justly assume that natural resistance may be largely a matter of inheritance. Whether this is to be interpreted as purely an instance of survival of the fittest or whether immunity

acquired by an individual can be wholly or in part transmitted to the offspring is an open question—at present in the same state of unclearness as are other questions relating to the transmissibility of acquired characteristics. However this may be, there are a number of facts available which indicate that inheritance plays an important part. It is apparent in the case of many diseases afflicting human beings that infection takes a milder course in those races among which it has long been endemic—whereas the same disease, suddenly introduced among a new people, is relatively more severe and spreads more rapidly. This seems to be the case with yellow fever and tuberculosis, and in measles and small-pox, too, the principle seems to hold good. Syphilis when first described authentically—as epidemically sweeping through Europe toward the close of the 15th century—appears to have been a far more acute and violent disease than it is among us to-day. It may well be that this depends upon a gradual elimination (elimination in this case, especially as far as reproduction is concerned) of those individuals that are fortuitously more susceptible and, by natural selection, a higher racial resistance is gradually developed. Whether or not direct inheritance of the individually acquired immunity can be considered at all as a contributing factor is difficult to decide. That immunity can be transmitted from mother to offspring was observed by Chauveau⁷ as early as 1888. Lambs thrown by anthrax-immune ewes possessed a higher resistance against this infection than did the lambs of normal ewes. The extensive experiments of Ehrlich,⁸ carried out chiefly upon mice with the vegetable poisons ricin and abrin, showed that in these cases immunity may be transmitted from mother to offspring, but depends upon a passive transfer of the specific antitoxins both by the blood and the milk of the mother. The sperm of the father did not seem to have anything to do with inherited resistance, since no immunity followed in the offspring when immunized males were paired with normal females. From the complete absence of immunity in the second generation (grandchildren) of the immunized female, and from the short duration (2 to 3 months) of its persistence, he concluded that the ovum itself had no influence, but that the entire phenomenon was attributable to a passive transference of antitoxins from mother to child during gestation and lactation. He interpreted, in the same sense, Chauveau's anthrax experiments, and similar experiments of Thomas⁹ and Kitasato¹⁰ with symptomatic anthrax, suggesting that, here also, a transept of antibodies from mother to offspring had taken place. The experiments of Ehrlich permit of no doubt as to the validity of his conclusions. However,

⁷ Chauveau. *Ann. Pasteur*, 1888.

⁸ Ehrlich. *Zeitschr. f. Hyg.*, 1892, Vol. 12.

⁹ Thomas. *Compt. rend. de l'acad. des sc.*, Vol. 94, cited by Ehrlich, *loc. cit.*

¹⁰ Kitasato. Cited by Ehrlich, *loc. cit.*

we must remember that they were carried out with antitoxic immunity only, in which the resistance is purely dependent upon the circulating antibody and is never, even in actively immunized individuals, a permanent state. In immunity such as that acquired against typhoid fever, plague, cholera, and other diseases after recovery from an attack, the individual remains relatively resistant long after the demonstrable antibodies have disappeared from the circulation, and we must assume that this permanent resistance depends upon a physiological alteration—inexplicable for the present, but surely residing in the body cells. In such cases it is by no means certain that there may not be a very slight, but through generations gradually accumulating, inheritance of immunity. At any rate the experiments of Ehrlich do not disprove such a possibility. Moreover, in this connection it must not be forgotten that natural immunity, unlike acquired immunity, cannot be passively transferred from one animal to another, and implies therefore a fundamental cellular difference rather than a condition depending merely upon antibodies circulating in the blood.

For this last reason also it has been unsatisfactory to attempt explanations of natural immunity purely upon grounds of bactericidal and other properties of the blood serum. These points we will take up at greater length when we discuss the mechanism of resistance in general.

An important observation upon the inheritance of serum properties is that which has been made by Ottenberg and Epstein¹¹ in connection with the iso-agglutinins. We shall see in another section that the blood serum of one human being will often possess the property of agglutinating the human blood cells of another individual. These iso-agglutinating constituents of the serum are apparently transmitted from parents to offspring. Von Dungern and Hirschfeld,¹² in studying these iso-agglutinins in 72 families, upon 348 people, not only confirmed the observations of the preceding workers, but showed that such inheritance follows Mendelian laws. Not only is this of great biological interest, but it is of great importance in connection with our present discussion in showing that such properties as agglutinating powers of serum can be influenced by inheritance from the father as well as from the mother.

The individual differences in resistance which unquestionably exist among members of the same species and races are very difficult to explain, but, as far as we can tell anything about them at all, they seem to depend upon variation in what is popularly spoken of as "general condition." The laboratory animals with which most experimentation is done, rabbits and guinea pigs, if healthy, show very

¹¹ Ottenberg and Epstein. *Proceedings of the N. Y. Path. Soc.*, 1908.

¹² Von Dungern and Hirschfeld. *Zeitschr. f. Immunitäts.*, Vol. 4, 1910.

slight individual variations. In fact, the astonishing uniformity of reaction on the part of guinea pigs of similar age and weight against measured quantities of bacterial toxins has alone made it possible to utilize these animals in the standardization of antitoxins. Pneumococcus and streptococcus cultures can be measured with reasonable accuracy upon white mice of approximately uniform weight, and the same animals are relatively uniform in their reactions to identical amounts of tetanus poison. Many other examples might be cited which make it clear that healthy animals of the same species, kept under the same conditions, fed upon the same food, and of approximately the same age and weight, differ but slightly from each other in reaction to the same infectious agent. This would indicate that the individual differences in resistance displayed so plainly by human beings are due, not to any *fundamental* individual variations, but rather to such fortuitous factors as nutrition, metabolic fluctuations, temporary physical depression, fatigue, or chilling. A person suffering from functional impairment of any kind is more likely to permit the invasion of a pathogenic micro-organism than is a perfectly healthy well-nourished individual of the same species.

Most of these facts we know from the accumulated experience of clinicians who also have given us much valuable information concerning the susceptibility to infection on the part of chronically diseased persons, especially diabetics and nephritics. In the case of a few of these influences, chilling and fatigue, experimental data on animals are available. It is, however, extremely difficult to analyze the causes underlying such depression of resistance. For instance, with fatigue or chilling there may be temporary congestion of mucous surfaces, due to vasomotor influences, which alter the secretions on mucous surfaces, or interfere with the normal mobilization of leukocytes, permitting penetration of bacteria where ordinarily they would have been held back. Our ignorance is nowhere more clearly illustrated than in the fact that we know practically nothing concerning the relation between a thorough chilling and the acquisition of what is spoken of as a common "cold." We can only assume that there is interference in some way with the normal bactericidal and phagocytic mechanisms, making possible the penetration and lodgment of small quantities of bacteria, ordinarily destroyed immediately after entrance or prevented from entering at all.

Of course we must except those individual differences of susceptibility which may be dependent upon inheritance. We know, for instance, that in such diseases as diphtheria, where resistance depends upon antitoxins circulating in the blood, there may be a passive immunity, conferred from mother to offspring, which lasts for several weeks or months after birth. It is important to remember such a possibility in the selection of guinea pigs for diphtheria antitoxin standardization, as Anderson has pointed out. Whether

or not a tendency to tuberculosis can be inherited is still an open question. In most cases it is more than probable that the supposedly inherited tendency to tuberculosis is not really an inherited susceptibility, but rather an actual infection acquired during childhood from the parents. Cornet and Kossel,¹³ who have recently summarized the statistics dealing with this problem, have come to the conclusion that this factor, namely, infection from the parents, probably is the cause of the greater frequency of tuberculosis among children of tuberculous parents, and that there is no definite proof of inherited susceptibility.

ACQUIRED IMMUNITY AND IMMUNIZATION

We have outlined in the preceding pages the differences in susceptibility to various diseases apparent among different species of animals, and have noted that the degree of resistance of some animals to infection with germs rapidly fatal to others is often sufficiently well-marked to be termed "immunity." Such immunity, because it is a natural biological attribute of the species, as much a characteristic property as are its anatomical or physiological properties, has been spoken of as "*Natural Immunity*."

It is a matter of common knowledge, however, that among species of animals readily susceptible to certain infections resistance, or even extreme resistance, i. e., immunity, may be acquired by an attack of the disease. Thus human beings who have recovered from plague, small-pox, typhoid fever, cholera, the exanthemata, mumps, typhus, yellow fever, and a number of other conditions do not ordinarily contract the disease a second time. In some of these conditions, notably cholera, plague, typhoid fever, and small-pox, the rule is almost invariable. In others, such as measles, scarlet fever, and mumps, a second attack may occur, though it is rare.

The following table briefly indicates infectious diseases in which permanent immunity follows an attack:

Infectious Diseases in Which One Attack Conveys Lasting Immunity

Plague.

Typhoid—second attack rare—about 2.4 per cent. (Curschmann).

Cholera.

Small-pox—second attack very rare.

Chicken-pox—second attack very rare.

Scarlet fever—second attack about 0.7 per cent.

Measles—second attack uncommon, but less rare than scarlatina.

Yellow fever.

Typhus fever.

Syphilis—reinfection rare, though more common than formerly supposed.

Mumps—second attack rare (Kraus).

¹³ Cornet and Kossel in "Kolle u. Wassermann," Vol. 5, 2d Ed.

No lasting immunity is conferred by one attack in:

Infection with the Pyogenic cocci
Gonorrhea
Pneumonia
Influenza
Glanders
Dengue fever
Diphtheria in general protection, second attack in 0.9
per cent. cases—0.01 antitoxin unit per c. c. of circu-
lating blood protects.
Recurrent fever
Tetanus
Erysipelas
Beri beri
Malaria
Tuberculosis

These observations actually form the point of departure of that entire branch of medical science which devotes itself to the study of resistance to infection, serum diagnosis, and specific therapy, and it will be seen that all the facts that have been gathered upon these subjects are the fruits of detailed analysis of this phenomenon of acquired immunity.

Its occurrence in many instances has been so striking that ancient observers, long before the birth of rational medicine, referred to it, and often drew from it conclusions of great hygienic importance. Thucydides, in the second book of his account of the Peloponnesian Wars, in describing the plague at Athens, notes the apparent safety from reinfection of those who had recovered, suggesting the possibility of their being therefore immune against disease in general. The literature of the Middle Ages and of earlier modern times contains numerous further references which indicate that acquired resistance was clinically recognized as a result of recovery from many diseases. The phenomenon was not only observed, but put to practical utilization by the ancients of China and India. Thus the practice of inoculating children with small-pox material from the active pustules of patients, or making them sleep in beds or wear the shirts of sufferers was a dangerous practice but logical, on the reasoning that the disease conveyed to a person in full health and good condition would probably take a mild course and confer immunity, while the naturally acquired disease, contracted often because of the weak and debilitated condition of the individual, would be more apt to end fatally.

Such methods, though barbaric and eventually unjustified by the naturally high mortality incident upon them, were actually brought to Europe from the East, and for a time practiced in European countries.

The first great advance which bridged the gap between the obser-

vations regarding naturally acquired immunity and rational experimental immunization was made by Edward Jenner. It had been noticed before Jenner began his work that milkmaids and others who had contracted cow-pox in the course of their occupations were usually spared when a small-pox epidemic occurred in their community. Sporadic attempts had been made to put this observation to practical use, but no one with sufficient intelligence, persistence, and training had taken up the matter seriously. Jenner, interested by the reports of this nature and by his own observations, was especially impressed by the similarity between the local manifestations of small-pox, cow-pox, and a disease of horses spoken of as "grease." Though at first disinclined to identify small-pox with cow-pox (at present the prevailing opinion is that the second is an attenuated form of the former), Jenner thoroughly investigated cases of alleged protection by cow-pox, a claim which before this had been hardly more than a rumor, and finally, with the encouragement of John Hunter, proceeded to the vaccination of human beings with cow-pox, testing the result by subsequent inoculation of the same individual with small-pox. His report to the Royal Society in 1796 and his subsequent publications incorporate the results of these experiments by means of which the practice of vaccination against small-pox was introduced, and the virtual eradication of the disease from civilized communities was attained.

The principles underlying small-pox vaccination are extremely simple. The attenuated virus after inoculation incites a mild and localized form of the disease, from which the subject recovers rapidly and completely. The recovery implies the mobilization of certain protective forces and a specific physiological alteration of the body in such a way that a permanently, or at least prolongedly, increased resistance against the disease remains. In consequence, if the individual is subsequently exposed to spontaneous infection with this disease, his acquired specific resistance suffices to prevent invasion by the virus. This is merely an artificial imitation of the conditions which obtain when an individual recovers from an attack of a disease and is rendered immune thereby. In this case, however, the attenuation of the virus has eliminated the dangers attendant upon an actual attack. The immunity thus conferred is probably never as perfect nor as lasting as that following a seizure of the disease in its unattenuated form; however, it suffices, as a rule, to prevent spontaneous infection which is never as severe a test as experimental inoculation.

In contrast to the "Natural Immunity" which is an inherited attribute of race or species, we speak of such increased resistance in a member of an originally susceptible race as "Acquired Immunity." When the immunity has been attained as the result of an attack of the disease itself it is spoken of as "Naturally or Spontaneously Ac-

quired Immunity." When produced by some form of treatment with the virus of the disease, altered in such a way that an actual attack is avoided, we speak of it as "Artificially Acquired Immunity."

The premises of Jenner's reasoning were valid as his experiments were convincing. But knowledge regarding infectious disease and its causation by living germs was not developed until almost one hundred years later, by the work chiefly of Pasteur. For this reason no direct continuation of Jenner's work appeared until Pasteur made his communication upon Chicken Cholera to the Parisian Academy of Medicine in 1880. Though his investigations differed entirely from those of Jenner both in method and the nature of the disease with which they dealt, Pasteur recognized the similarity of the fundamental principles underlying both discoveries.

His observations took origin in a purely accidental occurrence. Cultures of chicken cholera which had been allowed to stand without transplantation and under aerobic conditions for periods of several months were found to have diminished in virulence. Inoculated into chickens, they failed to kill, giving rise in many cases to localized lesions only. It occurred to Pasteur that inoculation with such an attenuated culture might protect against subsequent infection with fully virulent strains and, indeed, experimental investigation of this idea proved to be correct. He developed a method of "vaccination" against chicken cholera which consisted in injecting first a very much attenuated culture of the organism (*premier vaccin*), and, after 12 or 14 days, another less perfectly attenuated (*deuxième vaccin*), since he observed that a single inoculation was often insufficient to confer protection. After two inoculations a degree of immunity could be attained which sufficed to protect against spontaneous infection as well as against experimental inoculation with doses of the virulent germs, fatal for untreated animals.

These experiments, simple as they are, constitute the beginnings of the science of Immunity, since, for the first time, an investigator working with a pure culture of a pathogenic micro-organism had succeeded, in planned and purposeful experiments, in conferring artificial immunity. The path was now clearly indicated and the years immediately following were fruitful in the development of many methods by which pathogenic bacteria may be attenuated and changed in such a way that they can be used to confer immunity without causing more than a transient and harmless reaction in the subject. Most of the earlier discoveries of this kind came from Pasteur himself and from members of his school.

Since in all these methods the inoculated animal attains its increased resistance by reason of the activities of its own tissues, these processes are spoken of as "*Active Immunization.*" No protective factor is conferred directly. The disease itself is inoculated, though

in an altered form, and the subsequent immunity is purely the result of the physiological reaction occurring as the subject struggles against and overcomes the injected virus, bacteria, or their products. Such "Active Immunization," we shall see, is in contrast to "*Passive Immunization*," a procedure in which the serum of an actively immunized animal is injected into another, carrying with it certain substances by which protection is conferred. The recipient here is passively protected by products of the active reaction which has taken place in the body of the donor.

After his success in active immunization against chicken cholera Pasteur applied the principles here learned to experiments upon the protection of animals against anthrax. This problem was fraught with considerable difficulty because of the great virulence of the anthrax bacillus. However, successful attenuation was attained by a method which depended upon the cultivation of anthrax cultures at temperatures above the optimum for its growth. Toussaint¹⁴ had shown that the resistance of sheep could be increased if they were inoculated with blood from animals dead of anthrax after this had been heated to 55° C. for 10 minutes. Toussaint's idea had been that by heating the blood in this way the bacteria themselves were killed. Pasteur¹⁵ showed, however, that this was not the case, but that what actually occurred was a reduction of the virulence of the strain by the exposure to heat. As a matter of fact, moreover, the method of Toussaint did not furnish a reliable means of attenuating anthrax, and Pasteur succeeded in developing a far more satisfactory procedure on which he based a practical method for the protective vaccination of sheep and cattle.

His method was as follows:¹⁶ Virulent anthrax bacilli were cultivated at 42° to 43° C. on neutral chicken bouillon (Sobernheim states that horse or beef broth—or even agar—answers the same purpose). Cultivated under these conditions a gradual and progressive reduction of virulence occurs. After about 12 days of such cultivation the culture as a rule no longer kills rabbits, but is still virulent for guinea pigs and mice. After twenty-four or more days the virulence for rabbits and guinea pigs is lost and mice only can be killed with it. The latter—the most fully attenuated strain—was called *premier vaccin* by Pasteur, and, in the immunization of cattle or sheep, is first injected. After 10 or 12 days the stronger *deuxième vaccin* is administered. This is the method which Pasteur used in his now classical experiments at Pouilly-le-Fort, in which he convinced a hostile audience of the efficacy of his immuniza-

¹⁴ Toussaint. *Compt. rend. de l'acad. des sc.*, 1880.

¹⁵ Pasteur, Chamberland and Roux. *Compt. rend. de l'acad. des sc.*, Vol. 91, 1881.

¹⁶ Cited from Sobernheim. "Kraus und Levaditi Handbuch der Technik, etc.," Vol. 1, 1909.

tion. Sheep were protected in the manner indicated, and 14 days after the last injection a fully virulent culture was inoculated and the animals found capable of successfully resisting it.

In the train of this work many other methods of producing active immunity have been devised—all of them of considerable theoretical interest and many of them practically adapted to some special case. We may conveniently classify these methods as follows:

I. IMMUNIZATION WITH LIVING BUT ATTENUATED CULTURES

(1) Methods in which the attenuation is obtained by heating. This is the method of Toussaint as outlined above, in which anthrax blood was heated to 55° C. for 10 minutes, and is probably the least efficient or reliable method for the attenuation of the anthrax bacillus. It has been applied to rabies by Babes (cited from Kraus in "Kraus u. Levaditi Handbuch, etc.," Vol. 1, p. 708), who attenuated the virus by heating to 58° C. for periods varying from 2 to 40 minutes.

(2) Attenuation by prolonged cultivation of the bacteria at temperatures above the optimum for their growth. This is illustrated by Pasteur's anthrax immunization as described in the preceding paragraphs.

(3) Attenuation by passage through animals. Examples of this are Pasteur's experiments with the "rouget" organism, in which passage through rabbits diminished the virulence for hogs. The attenuation of rabic virus by passage through monkeys is another instance, and Jennerian vaccination is also an example of this, although here the attenuation by passage through cattle is attained naturally and not by experimental procedures. Based on the same principle is Behring's¹⁷ method¹⁸ of immunizing cattle against tuberculosis by inoculating them with tubercle bacilli of the human type.

(4) Attenuation by prolonged growth of bacteria on artificial media in the presence of their own metabolic products. This is the method first employed by Pasteur in chicken cholera, as described above, and is applicable to many organisms, such as pneumococci, streptococci, and others. In fact, it is difficult to maintain the virulence of many of these bacteria unless special methods of cultivation or passage through animals are practiced. Pasteur believed that free access of oxygen to the cultures increases the rapidity of the attenuation.

(5) Attenuation by drying. The classical example for this method is the Pasteur method of prophylactic immunization against

¹⁷ Behring. "Therapie der Gegenwart," April, 1907.

¹⁸ See also Römer, "Kraus u. Levaditi Handbuch," 1st Suppl., p. 310.

rabies. Rabbits are inoculated with *virus fixe*, and their spinal cords dried for varying periods in bottles containing KOH at a temperature of about 25° C. The virus grows progressively weaker with each day of drying. Greater details concerning this method are given in another place (see page 489).

(6) Attenuation by the use of chemicals.—Chamberland and Roux¹⁹ succeeded in attenuating anthrax by growing it in the presence of various antiseptics. They used carbolic acid 1 to 600, bichromate of potassium 1 to 1,500 and sulphuric acid 1 to 200, and found that, after a short time of cultivation under such conditions, the bacilli lost their ability to form spores and became avirulent for sheep. Behring²⁰ and others have applied this method to the attenuation of diphtheria toxin; Behring adds terchlorid of iodine, Roux potassium iodid—iodine solutions. The principle, of course, is not exactly the same in the last cases, since here the attenuation is not of the bacteria themselves, but rather of the toxin.

(7) Attenuation by cultivation under pressure. This method is difficult to apply, and has no striking advantages over other procedures. It was employed by Chauveau²¹ for the attenuation of anthrax. He succeeded in accomplishing this by cultivation of anthrax bacilli at 28-39° C. at a pressure of 8 atmospheres.

II. ACTIVE IMMUNIZATION WITH FULLY VIRULENT CULTURES IN SUBLETHAL AMOUNTS

The original methods of Pasteur carried out with chicken cholera and anthrax were aimed particularly at diminution of virulence, since these organisms, as isolated from the diseased animal, are so extremely infectious that it would be very difficult—(in the case of many animals, impossible)—to inoculate with the unattenuated germs without producing fatal disease. However, in the case of many other infections it has been found feasible to inoculate normal animals with the fully virulent germs in such small quantities that the body can successfully overcome them, and, in doing so, acquire specific resistance. It is obvious that this method is more easily carried out with the organisms which Bail terms “half parasites” than with organisms as highly infectious as anthrax. Ferran²² applied this method both to animals and to human beings with broth cultures of cholera spirilla. Högyes²³ has introduced a similar procedure for immunization against rabies by injecting dilutions of

¹⁹ Chamberland and Roux. *Compt. rend. de l'acad. des sc.*, 96, 1882.

²⁰ Behring and Wernicke. *Zeitschr. f. Hyg.*, 12, 1892.

²¹ Chauveau. *Compt. rend. de l'acad. des sc.*, Vol. 98, 1884.

²² Ferran. *Compt. rend. de l'acad. des sc.*, 1885.

²³ Högyes. “Lyssa Nothnagels Handbuch, etc.,” Vienna, 1897.

fully virulent rabie virus, beginning with a dilution of 1 to 10,000 and rapidly working up to a dilution of 1 to 10. In tuberculosis immunization with fully virulent cultures in small amounts has been attempted by Webb, Williams, and Barber,²⁴ using the Barber method of isolation, and giving a single micro-organism at the first injection. That such a method is feasible, if carried out with sufficient care, even with the most virulent germs, was demonstrated by the same workers. They succeeded in immunizing animals against anthrax (with cultures kept 12 hours on agar)²⁵ by injecting a single thread (3 to 6 bacilli) as the first dose, and then gradually increasing the amount.

In the general laboratory immunization of animals treatment with virulent bacteria in sublethal doses is of considerable value and frequently employed.

It would seem that possibly this method or some modification of it will be found to have very definite advantages over methods in which either attenuated or dead bacteria are employed. Bail's work upon the aggressins and upon anti-aggressin immunity (see chapter I, page 21) has opened the possibility that virulent bacteria provide, within the living body, specific aggressive substances which are not produced in the test tube. If this proves to be true, and the question is by no means settled, it may be necessary in such cases to immunize with organisms which are in a condition capable of producing these aggressins. Sublethal doses of fully virulent organisms would furnish these conditions more perfectly than attenuated avirulent strains, in which the invasive (aggressive) power is considerably diminished.

The methods of active immunization so far described differ from those which are to follow in that the preceding were all based upon the use of living bacteria or virus, whereas the methods to be described below depend upon the treatment of animals with dead bacteria or bacterial products. It is well to call attention in this place to the fact that a number of recent investigations seem to point to the greater efficiency of immunization with living germs. This method has recently given hopeful results in the case of plague in the hands of Strong;²⁶ and Metchnikoff and Besredka,²⁷ in their attempt to vaccinate chimpanzees against typhoid fever, make the statement that vaccination with dead typhoid bacilli or autolysates does not confer adequate protection, but that this can be attained by treatment with small doses of the living bacilli.

²⁴ Webb, Williams, and Barber. *Jour. of Med. Res.*, Vol. 15, 1909.

²⁵ This was not possible where the organisms were taken directly from the blood of a dead mouse. In such cases even a single thread caused fatal disease.

²⁶ Strong. *Jour. of Med. Res.*, May, 1908.

²⁷ Metchnikoff and Besredka. *Ann. Past.*, Vol. 25, 1911.

In speaking of this subject it is well to mention recent observations upon immunization with "*sensitized*" bacteria,²⁸ although this necessitates anticipatory reference to subjects not so far discussed. It is a matter of common experience in laboratories that rabbits and other animals will withstand relatively large amounts of pathogenic bacteria if these are first treated with heated specific immune serum (sensitized). This is probably due to the fact that such "sensitized" micro-organisms are very rapidly taken up by phagocytes. In spite of the phagocytosis, immunity is developed. Metchnikoff and Besredka, in the communication alluded to above, state that typhoid vaccination with unaltered living bacilli is efficient, but is attended by severe local and general reactions. If the living bacilli are first "sensitized" no such severe reaction occurs and immunization is nevertheless successful. The recent work of Gay points in the same direction, and it is at least possible that by the practice of sensitization we may be able to employ living unattenuated organisms for purposes of immunization more extensively than we have in the past.

III. ACTIVE IMMUNIZATION WITH DEAD BACTERIA, AND BACTERIAL EXTRACTS

This method is the one most extensively practiced in the laboratory immunization of animals. It is usual in most experiments of this kind to inject dead organisms once or twice before living bacteria are administered. High degrees of resistance can in some instances be attained by progressively increasing doses of dead cultures only. This method is not only useful in experimental work, but is clinically employed in the active immunization of human beings as introduced by Wright and as applied, before Wright, to tuberculosis (tuberculin treatment). But it is very probable that the immunity so attained is not entirely comparable to the immunity following an attack of a disease, nor even that produced by the injection of living bacteria.

The method employed for killing the bacteria is of considerable importance since, both by excessive heating as well as by too vigorous chemical treatment, the immunizing properties of the bacterial protein may be destroyed. In employing heat it is a safe rule never to expose the bacteria for prolonged periods to temperatures which considerably exceed the thermal death point. As a rule, heating non-spore-forming bacteria to a temperature of from 65° to 70° C. for

²⁸ Refer to p. 159 and the discussion of the conception of "sensitization" which follows.

thirty minutes will suffice to kill them without too radically altering the immunizing properties of the protein constituents.²⁹

If the temperature is not raised above 60° C., and this is advised by many workers, the suspensions must be carefully controlled by cultural tests before they are used, at least for the treatment of human beings. As we shall see in a later section, the best results have been obtained when heating was not carried beyond 53° to 55° C.

When bacterial death is to be accomplished by chemicals the antiseptics most commonly used are carbolic acid (0.5 per cent.), toluol (removed before use of vaccine by filtration or evaporation), chloroform, and formaldehyd (1 per cent.).

Pfeiffer, who was one of the first to practice the immunization of animals with dead bacteria on an extensive scale, believed that, in the case of bacteria which were toxic by reason of their intracellular constituents (endotoxins), the injection of the cell protein itself, whether dead or alive, was the sole essential for successful immunization. The method developed by Kolle³⁰ and by Pfeiffer and Marx³¹ for the prophylactic immunization of human beings against cholera depends upon the injection of cholera cultures emulsified in salt solution, killed by exposure to 58° C. for one hour, and further insured against contamination by the addition of 0.5 per cent. phenol. The application of this method to other diseases, both prophylactically and therapeutically, is more fully discussed in another place. (See chapter XIX.)

Since the essential point in such immunization is the introduction of the bacterial protein, it is often customary to inject bacterial extracts instead of the whole cells. This has been especially desirable in the case of such insoluble micro-organisms as the tubercle bacillus, where the injection of the whole dead organism produces localized reactions similar to those caused by the living bacteria.³² Thus "Old Tuberculin," as commonly used, is a glycerin-broth extract of tubercle bacilli. The method has been extensively used and a variety of procedures have been devised for bacterial extraction. These have included simple autolysis of the bacterial bodies in alkaline broth, shaking in salt solution in mechanical shakers, trituration with salt or sand, trituration after freezing, digestion with proteolytic enzymes, and extraction by pressure in a Buchner press.

We may mention some of the more important methods for pre-

²⁹ In a subsequent chapter (p. 258) we shall see that the physical changes produced in an antigen by heat result in differences in the antibodies formed after animal inoculation. This point has practical significance in the present connection. See also the chapter on agglutinins, the work of Joos there discussed, and Friedberger and Moreschi, *Centralbl. f. Bakt.*, 1905, Vol. 39.

³⁰ Kolle. *Deutsche med. Woch.*, 1897, p. 4.

³¹ Pfeiffer and Marx. *Deutsche med. Woch.*, 1898.

³² Prudden and Hodenpyl. *N. Y. Med. Journal*, 1891.

paring bacterial extracts for purposes of immunization and antigen production in general as follows:

A. Extraction of Bacteria by Permitting Them to Remain for Prolonged Periods in Fluid Media

The bacteria may be grown upon slightly alkaline bouillon and kept at incubator temperature for one to two months. They are then filtered through Berkefeldt or other suitable filters. This is the common method of producing antigen for precipitin reactions, in fact the method employed by Kraus in the discovery of the bacterial precipitins. It is by no means certain whether the antigens prepared in this way represent simple extractions or autolytic products of the bacteria; probably both processes take place. The antigenic value of the fluids obtained in this way is never very great. From such filtrates Brieger and Mayer, Pick, and others have attempted to obtain the antigen in a purified form by chemical precipitation. Pick³³ precipitates the bouillon filtrate by saturation with ammonium sulphate; the precipitate is redissolved in water and again precipitated with ammonium sulphate and the resultant precipitate dried on a filter. It is then dissolved in water and precipitated with alcohol. The sticky substance which comes down represents the antigen.

Suitable extracts can occasionally be obtained also by emulsifying agar cultures in physiological salt solution and allowing them to stand for twenty-four hours or more at incubator temperature. In our own experience we have found this method rather inefficient for yielding strong extracts. More efficient extraction is usually obtained when the bacteria are suspended in alkaline fluids such as $\frac{N}{10}$ sodium hydrate. Lustig and Galleotti digest the bacterial mass for 24 hours with 1 per cent. NaOH, then precipitate with ammonium sulphate, dry in vacuo and pulverize.³⁴

Recently, also, Uhlenhuth³⁵ has employed the proprietary preparation "antiformin"³⁶ for the production of antigens. This

³³ Pick. "Hoffmeister's Beiträge, etc.," Vol. 1, 1902. For an extensive discussion of the various methods employed for the production of bacterial antigens by chemical methods see Pick in Kraus und Levaditi, etc., Vol. 1, and the same author in Kolle u. Wassermann, etc., 2nd Ed., Vol. 1.

³⁴ See Pick. *Loc. cit.*

³⁵ Uhlenhuth. *Centralbl. f. Bakt.*, I, Ref. Vol. 42, Beilage, p. 62.

³⁶ "Antiformin" is a substance largely employed for the cleansing of pipes and vats of organic matter because of its powerfully solvent action. Its value in concentrating tubercle bacilli out of sputum and other mixtures depends upon its power to dissolve the tissue elements and all bacteria except those that are acid-fast. Rosenau ("Preventive Medicine and Hygiene,"

substance thoroughly dissolves all but the acid-fast bacteria when used in concentrations of 2.5 per cent. Since it is alkaline it is necessary to neutralize it with hydrochloric or sulphuric acid before use.

For the preparation of antigen from pneumococci Neufeld³⁷ has utilized the solvent action upon these organisms of bile. He adds the bile and broth cultures just as this is done in the diagnostic "bile test" (0.1-0.2 c. c. of fresh bile to a broth culture; sodium taurocholate solution can also be used). Many bacteria can also be broken up by emulsifying them in 17 per cent. salt solution and allowing them to stand for some time in this medium and then diluting this to 0.85 per cent.

B. Extraction by Mechanical Methods

One of the most useful methods for obtaining extracts of bacteria within a relatively short time is that which Besredka³⁸ has applied mainly for the preparation of typhoid (endotoxin), 24-hour agar cultures washed up in very small quantities of physiological salt solution, killed by heat at 60-65° C. and dried *in vacuo*. The dried mass is mixed with a measured quantity of dry salt and the mixture thoroughly triturated in a mortar for a considerable time. While triturating distilled water is added in small quantities until the fluid represents a 0.85 per cent. salt solution. This is allowed to stand for anywhere from a few hours to a week, and the bacteria are then removed by centrifugalization. This method has been modified by many observers and gives good results whenever thorough trituration is practiced. It is also probable that the exposure to the hypertonic salt solution in the earlier stages of the trituration may aid considerably in breaking up the bacteria.

Trituration after freezing is a method which has yielded excellent results in the hands of Macfadyen and others. This requires a rather complicated piece of machinery originally described by Macfadyen and Rowland. The principle of this is one of mechanical trituration in a steel cylinder which is surrounded by an ice-brine mixture so that the bacteria and sand may be kept frozen during the process.

Appleton, 1913, p. 1020) gives its composition as follows: "Antiformin consists of equal parts of liquor sodæ chlorinate of the British Pharmacopœia and a 15 per cent. solution of caustic soda. The formula for liquor sodæ chlorinate is as follows:

| | |
|------------------------|--------|
| Sodium carbonate | 600 |
| Chlorinated lime | 400 |
| Distilled water | 4,000" |

³⁷ Neufeld. *Zeitschr. f. Hyg.*, Vol. 34, 1900.

³⁸ Besredka. *Ann. de l'Inst. Past.*, 19-20, 1905, 1906.

Mechanical trituration is also the principle of the production of the new tuberculins as advised by Koch.

One of the earliest methods of obtaining bacterial substances by mechanical means was that used by Buchner and Hahn³⁹ in the production of their "plasmines." The bacteria were grown in quantity on large agar surfaces, the moist bacterial masses trituated together with quartz and were then subjected to high pressure in an especially constructed press spoken of as the "Buchner press."

Mechanical breaking up and extraction of the bacteria also underlies in principle the use of the variously constructed shaking machines. There are many models of such machines on the market, all of them designed to accomplish prolonged agitation of bacterial emulsions. In many cases the apparatus can be placed inside of an incubator and shaking carried on at 37.5° C. The bacteria are suspended for this purpose in distilled water salt solution, weak alkali, or in serum, and glass beads or sand may be added to aid in their mechanical injury. Shaking must be continued for 24 hours or more in order to give good results.

IV. ACTIVE IMMUNIZATION WITH BACTERIAL PRODUCTS (TOXINS)

As soon as the investigations of Roux and Yersin had shown that in some diseases, at least, the injury sustained by the infected animal was largely due to the soluble toxins produced by the bacteria, it was logical to attempt to immunize animals with such products. Probably the first attempts in this direction were those made by Salmon and Smith in hog cholera. The experiments of these writers have attained much historical importance since they represent the first purposeful attempt to immunize animals with the products of bacterial metabolism. In the actual experiment, however, the immunization practiced by Salmon and Smith was probably a combination of immunization by bacterial products and by dead bacteria. Nevertheless, the thought of immunization with bacterial products was the underlying one in their experiments. Working with the hog cholera bacillus which they had recently discovered they immunized pigeons in the following way: The bacilli were grown in broth for two weeks, and the cultures were killed by exposure to 58° to 60° C. for several hours. One and one-fifth cubic centimeter of this culture liquid was then injected into pigeons, and after three such injections the inoculated pigeons withstood, without harm, doses of the bacilli which rapidly killed untreated animals. Salmon and Smith⁴⁰ stated distinctly in their conclusions that: "Immunity may

³⁹ Buchner and Hahn. *Münch. med. Woch.*, 1897.

⁴⁰ Salmon and Smith on "A New Method of Producing Immunity from Contagious Disease," *Proc. Biol. Soc.*, Wash., D. C., III, 1884, 6, p. 29, printed Feb. 22, 1886.

be produced by introducing into the animal body such chemical products (results of bacterial growth in culture fluids) that have been produced in the laboratory.⁴¹

Similar attempts to immunize rabbits against certain forms of septicemia by the injection of culture filtrates were made by Chamberland and Roux⁴² in 1888,⁴³ and the same investigators applied this method to anthrax immunization just prior to the discovery of diphtheria toxin by Roux. However, neither in hog⁴⁴ cholera⁴⁵ nor in the other infections upon which this method was first tried do the bacteria produce a true soluble toxin, and the immunization which was accomplished depended probably upon the injection of bacterial extracts. Nevertheless, these attempts had shown the way in a new direction, and bore immediate fruit in the investigations of Brieger and Fraenkel,⁴⁶ and more especially in those of Behring^{47 48 49} and his collaborators. Fraenkel, though following the method of injecting filtered diphtheria culture fluids, came to the erroneous conclusion that the toxin and the immunizing substances in the cultures were not identical (*loc. cit.*, p. 1135).⁵⁰ The degree of immunity obtained in his experiments, moreover, was a slight one only.

Behring, in his first work, in collaboration with Kitasato, succeeded in immunizing animals with culture filtrates and with pleural exudates of diphtheritic animals. Similar results were accomplished with tetanus. Since the publication of these results—especially in consequence of the epoch-making discovery of passive immunization, to which they were the immediate guides, toxin immunization has been investigated and accomplished in all cases in which a true soluble toxin can be demonstrated. It has accordingly been carried out with the exotoxins of pyocyaneus⁵¹ bacilli, the bacilli of symptomatic anthrax⁵² and botulinus,⁵³ the specific leukocidins⁵⁴

⁴¹ For a copy of the original paper by Salmon and Smith I am indebted to Professor Theobald Smith.

⁴² Chamberland and Roux. *Ann. Past.*, Vol. 1, 1888.

⁴³ *Op. Cit.*, Vol. 2, 1889.

⁴⁴ Joest in "Kolle u. Wassermann Handbuch, etc.," Vol. 3, p. 632.

⁴⁵ Karlinski. *Zeitschr. f. Hyg.*, Vol. 28, 1898.

⁴⁶ Brieger and Fraenkel. *Berl. kl. Woch.*, 1890, Nos. 11, 12 and 49.

⁴⁷ Behring and Kitasato. *Deutsche med. Woch.*, No. 49, 1890.

⁴⁸ Behring. *Deutsche med. Woch.*, No. 50, 1890.

⁴⁹ Behring and Wernicke. *Zeitschr. f. Hyg.*, 1892.

⁵⁰ De Schweinitz, indeed, who further studied hog cholera immunization (*Medic. News*, 1892; *Centralbl. f. Bakt.*, Vol. 20, 1896), claimed that in sterilized milk the bacillus produced "enzymes" with which immunization could be accomplished.

⁵¹ Wassermann. *Zeitschr. f. Hyg.*, Vol. 22.

⁵² Kempner. *Zeitschr. f. Hyg.*, Vol. 23, 1897.

⁵³ Grassberger and Shattenfroh. *Deuticke*, Wien, 1904.

⁵⁴ Denys and Van der Veldé. "La Cellule," Vol. 2, 1895.

produced by staphylococci, and with various bacterial hemolytic poisons (tetanolyisin and other bacterial hemotoxins). The result of all this work has been the very important determination that susceptible animals may be actively immunized both against the effects of the toxin alone, as well as against the virulent bacteria themselves, by systematic treatment with culture filtrates containing the toxins. Since in many cases the effects of the toxins were so powerful that their attenuation was desirable, Behring and others have advised the addition of iodinterchlorid and other chemicals to the first injections.

PASSIVE IMMUNIZATION

In the logical development of the fundamental facts regarding immunization, with attention focused early on the blood and body fluids as the probable carriers of immunity, it was but a rational step from active immunization to the conception that such acquired immunity might be transferred from a treated to a normal animal by injecting blood from the former into the latter. This was probably the underlying thought of Toussaint's⁵⁵ early work with anthrax, in which he heated anthrax blood to 55° C. and injected it into other animals, wrongly believing that the bacteria had been killed by the heating. The method of Toussaint, however, was vague in its conception, and in no way constitutes an example of true passive immunization. The beginning was made in a purposeful and clearly conceived way by Richet and Héricourt.⁵⁶

These investigators actively immunized dogs against staphylococci, and then attempted to transfer the immunity to normal rabbits by injecting defibrinated blood from the immune dogs. Their success was a partial one only, for reasons that we will discuss directly. Reasoning similar to that of Richet and Héricourt was applied by Babes and Lepp⁵⁷ to rabies immunization. When the blood of rabies-immune dogs was injected into normal dogs and rabbits, and these inoculated with rabies several days later, the treated animals regularly survived the controls, but in one dog only was the occurrence of rabies absolutely prevented. Since their animals were not experimentally inoculated, but subjected to the more uncertain method of allowing them to be bitten by a mad dog, and since the series included 4 animals only (2 treated and 2 controls), Babes and Lepp were unable to draw definite conclusions. The establishment of passive immunization as a proved scientific fact was finally accom-

⁵⁵ Toussaint. *Compt. rend. de l'acad. des sc.*, 1880.

⁵⁶ Richet and Héricourt. *Compt. rend. de l'acad. des sc.*, 1888, Vol. 107, p. 750.

⁵⁷ Babes and Lepp. *Ann. Past.*, Vol. 3, 1889.

published in 1890-1892 by Behring and Kitasato,⁵⁸ and by Behring and Wernicke. The results of this work—the direct outcome of their success in actively immunizing with soluble toxins, is summarized in their first paper as follows: “The blood of tetanus-immune rabbits possesses tetanus-poison-destroying properties; these properties are demonstrable in the extravascular blood and in the serum obtained from this; these properties are of so lasting a nature that they remain active in the bodies of other animals, so that one is enabled to obtain positive therapeutic results by transfusing the blood or injecting the serum. These tetanus-poison-destroying properties are absent from the blood of non-immune animals, and when the tetanus poison is inoculated into normal animals it can be demonstrated as such in the blood and other fluids of these animals after death.”

With these researches begins the therapeutically practicable method of passive immunization which is now in such widespread and successful use in the treatment of diphtheria, in the prophylactic treatment of tetanus, and, to a less common and less successful degree, in the treatment of dysentery, typhoid fever (Besredka), plague, and a number of other bacterial diseases. The same method has been successful in the treatment of various diseases of domestic animals. The principle was also applied by Ehrlich⁵⁹ to ricin and croton immunity, in the formulation of which he succeeded in working out passive immunization on a quantitative basis, showing that the degree of immunity in such cases could be directly referred to the amounts of the specific antitoxin present in the blood of the immunized animal. Calmette,⁶⁰ and Physalix and Bertrand,⁶¹ then succeeded in producing passive immunization against snake venoms.

To summarize the success of passive immunization in general we may say that it has achieved its greatest usefulness in the case of those diseases in which the pathogenesis depends upon a true exotoxin—which, as we have mentioned before, leads to the formation of an antitoxin in the immunized animal. In these cases the passive immunization is accomplished by the transfer of the antitoxins from the treated to the normal animal.

In the case of bacterial infections in which no true toxin is formed—where no antitoxin results and the immunity depends, as we shall see, upon an enhancement of the bactericidal and phagocytic properties of the blood and the cells, passive immunization has not been a practical therapeutic success. The probable reasons for this cannot be properly discussed until we have examined more closely into the mechanism by which the immune animal is protected after

⁵⁸ Behring and Kitasato. *Deutsche med. Woch.*, No. 49, 1890.

⁵⁹ Ehrlich. *Deutsche med. Woch.*, 1891; *Fortschr. d. Med.*, p. 41, 1897.

⁶⁰ Calmette. *Compt. rend. de la soc. de biol.*, 1894.

⁶¹ Physalix and Bertrand. *Compt. rend. de la soc. de biol.*, 1894.

specific treatment with bacteria or their products. The moderately beneficial effects of the various antiplague sera and the limited success attending the use of antistaphylococcus, antistreptococcus, and antipneumococcus sera probably depend, as recent work tends to show, not upon the direct action of antitoxic bodies, but rather upon the indirect opsonic action^{62 63 64 65} which renders the bacteria more easily amenable to phagocytic action. These points we shall discuss at greater length in a succeeding chapter.

SPECIFICITY

In speaking of methods of immunization in the preceding sections we have frequently employed the terms "specific" and "specificity," without sufficiently defining them. It will be necessary to explain them since the principle of specificity is at the same time one of the most important and one of the most mysterious of the phenomena of immunity. When an individual has recovered from an attack of typhoid he is thereafter immune to typhoid—but to no other disease—similarly with plague, cholera, small-pox, etc. The same principle governs artificial immunization. Vaccination against anthrax protects against anthrax only—and active or passive immunization in any of the infectious diseases produces a resistance which is "specifically" aimed only at the particular infectious agent with which the original active immunity was produced. The principle points to an exquisite chemical difference between the protein substances which constitute the bacterial cell bodies or their metabolic products. For although by chemical methods we can detect no differences between them—yet the reactions of immunity are sharply differentiating. When we come to consider the antibodies which specifically precipitate the substances by which they are incited we shall see that the delicacy and consequent differential value of these reactions far outstrip any known chemical methods, and it is upon this principle indeed—inexplicable as it is—that the great diagnostic value which these reactions have attained depends. The conception of the specificity of the causes of infectious disease, as well as that of the specificity of toxins, has become so common and self-evident to us that we are too apt to forget how fundamental to progress the establishment of this fact was in the early days of bacteriological research. When, in 1878, Koch published his treatise on the "Etiology of Wound Infections" specificity was not generally accepted, and the supposed metamorphosis of bacterial species, as asserted by

⁶² Neufeld. *Deutsche med. Woch.*, No. 11, 1897.

⁶³ Neufeld and Rimpau. *Deutsche med. Woch.*, No. 40, 1904.

⁶⁴ Bail and Kleinhans. *Zeitschr. f. Imm.*, Vol. 12, 1912.

⁶⁵ Weil. *Zeitschr. f. Hyg.*, 75, 1913.

Hallier and others,⁶⁶ had first to be scientifically refuted by Cohn, Koch, and their pupils, before it could be assumed that a given infectious disease was always the result of infection with a definite and constant species of bacteria. The same applied to the specificity of toxins—and rational investigations into the reaction of the animal body against bacterial poisons was not possible until the works of Roux and Yersin on diphtheria and that of Kitasato on tetanus had differentiated between the true, specific bacterial poisons and the unspecific ptomains and “sepsins” of Selmi, Nencki, and Brieger.

⁶⁶ Hallier. Cited from Behring, “Bekämpfung der Infektionskrankheiten,” Leipzig, 1894.

CHAPTER IV

THE MECHANISM OF NATURAL IMMUNITY AND THE PHENOMENA FOLLOWING UPON ACTIVE IMMUNIZATION

ANTIBODIES AND ANTIGENS. THE ORIGIN OF ANTIBODIES

THE MECHANISM OF NATURAL IMMUNITY

PASTEUR'S work on active immunization was carried out in the later seventies and the early eighties. During and immediately after this time it was very natural that the attention of investigators should have concentrated upon the elucidation of the causes underlying both the natural resistance against bacteria observed in animals and man, and the changes which during active immunization were fundamentally responsible for the acquisition of resistance.

It was easily determined that there were no anatomically and physiologically determinable differences between the various mammalia which could account for the observed striking variations of susceptibility, nor could gross anatomical or histological changes be noted in an animal which had been artificially immunized. Morphologically such an animal was indistinguishable both in the size and appearance of its organs, and in the arrangement and structure of its cells from any other individual of the same species not subjected to treatment.

It was a natural development of the investigations brought to bear upon this problem that attention should, for a time, be concentrated upon the phenomena of inflammation, processes which were regularly associated with infections of all kinds and seemed indeed to represent a sort of local expression of tissue resistance to the invading micro-organisms.

It was in the course of investigations upon the nature of inflammation that Metchnikoff first became interested in problems of resistance. In 1883 he presented a paper at the Naturalists' Congress at Odessa, in which he referred the absorption of dead or foreign corpuscular elements in the bodies of invertebrates to a process of intracellular digestion carried out by phagocytic cells. As early as 1874 Panum had suggested that possibly resistance against invading micro-organisms might be due to a similar intracellular destruction,

and Metchnikoff, soon after his first communication, extended his phagocytic studies to phenomena of infection. His first investigations concerned themselves with an infectious disease caused by a form of yeast in a small crustacean—the daphnia or water flea. He showed that recovery or death from the disease depended upon the completeness with which the invading micro-organisms were taken up by the white blood cells found in the body cavity of the daphnia. Immediately subsequent studies, carried out with the aid of numerous pupils, embraced an extensive material throughout the animal kingdom in which he attempted to show parallelism between natural immunity and the phagocytic activities mobilized by the body against the invading germs.

Meanwhile studies along another path were in progress. It had been observed many years before this by the physician, Hunter, that the shed blood of animals was not as easily subject to putrefactive change as were many other organic substances. Similar observations by Traube and, in 1881, by Lord Lister¹ (the latter reported at a time when Pasteur's experiments were reaping their first practical results) further stimulated investigation of the blood as the possible seat of the increased antibacterial property. For, indeed, these observations seemed to imply that by resisting decomposition, even when inoculated with putrefying material, the blood must possess definite means of inhibiting or even destroying the putrefactive bacteria.

In 1884, in a dissertation submitted at Dorpat, Grohman² stated that cell-free blood plasma inhibited the growth of micro-organisms. But Grohman was unable to determine actual bacterial destruction. Similar, but inconclusive, observations were published by Von Fodor³ in 1887. In 1888, however, Nuttall,⁴ who was investigating the validity of the phagocytic theory of Metchnikoff, experimentally determined that normal blood possessed the property of killing bacteria—a property now spoken of as “bactericidal” power. The attitude taken by Nuttall, and others of the Flügge school, toward Metchnikoff's opinions was one of doubt as to the fundamental significance of phagocytosis in determining resistance. They argued that Metchnikoff had not yet proved that *living* bacteria were taken up by the phagocytic cell, and that the action of these cells might therefore be interpreted as merely a process of removal of the dead bacteria, after these had been killed by other influences. Nuttall, accordingly, repeated some of Metchnikoff's experiments on anthrax in frogs and rabbits, essentially confirmed the basic observations, but showed also that the cell-free defibrinated blood of these and

¹ Lister. *Trans. Intern. Med. Congress*, London, 1881.

² Grohman. Cited from Lubarsch, *Centralbl. f. Bakt.*, 6, 1889.

³ Fodor. *Deutsche med. Woch.*, No. 34, 1887.

⁴ Nuttall. *Zeitschr. f. Hyg.*, Vol. 4, 1888.

other animals possessed definite bacteria-destroying properties (bactericidal power) for many different micro-organisms. He detected similar properties in pleural exudates, pericardial fluids, and aqueous humor, and determined that this property was "inactivated" or destroyed when the fluids were heated to 55° C. for 10 minutes or longer. Buchner⁵ then confirmed Nuttall's results and showed further that the bactericidal property resided, not only in defibrinated blood, peptone blood, and plasma, but was present also in the serum obtained after clotting. He applied the term "*Alexin*" to this active constituent of the blood—likening its action to that of a ferment.

The immediate theoretical result of these discoveries was an attempt, begun by Flügge's school, to base natural as well as acquired resistance upon the bactericidal properties of the blood and body fluids in general. For the observations of Nuttall and Buchner were soon extended to peritoneal and other exudates by Stern,⁶ and to ascitic fluids by Prudden.⁷ By these two groups, that of Flügge-Nuttall-Buchner on the one hand, and that of Metchnikoff on the other, there were founded the two schools of immunity—the humoral and the cellular, both originating in attempts to explain natural immunity, and later extending to problems of acquired resistance. And it is to the diligent and ingenious intellectual and experimental conflict between these schools that we owe much of the knowledge we now possess concerning the phenomena of immunity. A bridge between them was early established when Buchner himself—even before Metchnikoff—suggested the possible leukocytic origin of the bactericidal serum constituent (alexin). The later work of Denys, of Gruber and Futaki, of Wright, of Neufeld, of Bail, and of others has demonstrated, as was to be expected, the inadequacy of either point of view by itself, and the intimate interdependence of the humoral and the cellular processes.

As concerns the relation of bactericidal serum effects and natural immunity, it could be unquestionably shown by Nuttall, Buchner, Nissen,⁸ and their immediate followers that the blood of most animals possessed bactericidal properties against many micro-organisms, their experiments being so planned that the participation of leukocytes could be absolutely excluded. However, a parallelism between bactericidal power and the degree of natural resistance could not be established. Lubarsch,⁹ writing during the early periods of the controversy, stated that "he would regard the (purely humoral)"¹⁰

⁵ Buchner. *Centralbl. f. Bakt.*, 1889.

⁶ Stern. *Zeitschr. f. klin. Med.*, Vol. 18 (cited from Hahn).

⁷ Prudden. *Med. Rec.*, Jan., 1890.

⁸ Nissen. *Zeitschr. f. Hyg.*, Vol. 6.

⁹ Lubarsch. *Centralbl. f. Bakt.*, Vol. 6, 1889.

¹⁰ Bracketed phrase our own.

experiments of Nuttall as decisively contradicting the phagocytic theory if the bactericidal action of the blood (for anthrax bacilli) could be shown to be more potent in immune than in susceptible animals." Metchnikoff¹¹ himself, taking this point of view, called attention to the fact that the blood serum of rabbits, animals that are highly susceptible to anthrax, is more powerfully bactericidal for these micro-organisms than is the blood of dogs or even that of immunized calves, both of which are much more resistant than are rabbits. Nuttall answered this by reporting that the blood of anthrax-immunized calves is actually more powerfully bactericidal than is that of normal calves. Although this argument of Nuttall was perfectly valid in principle, it exerted little influence on opinions at this time because anthrax happens as a matter of fact to belong to that group of infections in which bactericidal protection is actually secondary to phagocytic, and Lubarsch could show that the differences observed by Nuttall were often less than those obtaining between specimens of blood taken from individual normal rabbits.

Lubarsch himself, then, in carefully planned experiments, showed that rabbits and cats could be killed with quantities of anthrax bacilli far less than the number which the extravascular blood of these animals can destroy. He concluded that the resistance, in these cases at least, is certainly not parallel with the bactericidal properties of the blood, and suggested the possibility that the intravascular blood does not possess bactericidal power to the same degree in which it is possessed by the extravascular plasma or serum. This point, first raised by Lubarsch—namely, the possibility of a difference between the intravascular blood and the extravascular blood serum or plasma in bactericidal functions—soon became one of the focal points of the controversy, since Metchnikoff, admitting the bactericidal power of the shed blood, assumed that this was purely the result of substances given off by the leukocytic cell-bodies after extravascular injury.

The Metchnikoff school defended its premise by the dual method of attempting on the one hand to establish a parallelism between phagocytic activity and natural resistance, and, on the other hand, by showing that the cell-free blood serum of naturally resistant animals often furnished an excellent culture medium for the bacteria in question. Thus Wagner showed that anthrax bacilli grow well in the blood of fowls at 42° C., and Metchnikoff himself called attention to the fact that pigeons' blood is an excellent medium for the cultivation of the Pfeiffer bacillus, whereas the living pigeon is entirely insusceptible to influenzal infection. Arguments based on such observations, however, have lost much of their original weight, for we have since then learned more about the delicate quantitative

¹¹ Metchnikoff. *Virch. Archiv*, Vol. 97, 1884.

conditions and the difficulties of accurate measurements obtaining in experiments upon *in vitro* bactericidal phenomena. For although a specimen of the blood of a naturally immune animal may be capable of destroying a considerable number of bacteria of a given species, the implantation of such a specimen with a slight excess of the bacteria would soon exhaust the active serum constituents and profuse growth could then take place. Furthermore, the conditions of temperature established in cultural experiments lead rapidly to a deterioration of the alexin necessary for bactericidal action, and any bacteria remaining alive at the end of a number of hours would then have unopposed opportunity to multiply.

The attempts to establish parallelism between phagocytic activity and natural immunity, though somewhat more successful than the analogous efforts of the humoral school, nevertheless also failed to furnish complete explanation for existing conditions, and, as we shall see, no adequate generalizations could be made until later years revealed the close coöperation between cells and fluids. We must postpone any attempts to do justice to this phase of the problem, therefore, until we are in a position to discuss the question of phagocytosis on the basis of a fuller knowledge of the phenomena which influence it.

The clear thinking and unprejudiced logic brought to bear upon this controversy by some of the great bacteriologists of this time are nowhere more instructively illustrated than in a short introduction published by v. Behring¹² to his second article on diphtheria. He says: "Neither deduction nor theorizing can at present decide whether a compromise will be found in the future between the two hypotheses (humoral and cellular), or whether the one or the other alone will be found correct. As yet the opinions of many experimenting bacteriologists are in direct opposition in this respect. Meanwhile, for the purposes of medical advancement and therapeutic success it is not necessary to await a decision of this question. . . . It is indeed of advantage to the cause if the struggle against infection is undertaken from the most varied points of view; attempts to make proselytes for a dogma have never led to progress. In this sense I will try to summarize those experimental results which support the humoral point of view without attempting particularly to detract from the importance of opinions which I do not share."

THE PHENOMENA FOLLOWING UPON ACTIVE IMMUNIZATION

The cellular and humoral points of view, formulated largely upon the facts of natural immunity, were equally applied, almost from the beginning, to the explanation of active immunization. The light

¹² v. Behring. *Zeitschr. f. Hyg.*, Vol. 12, 1892.

thrown upon these phenomena by the efforts of both schools rapidly led to a complete abandonment of those earlier theories of immunity which had conceived the acquired resistance of animals against bacteria as a purely passive development in the body of conditions unfavorable for bacterial growth.

Among these earlier theories, now of historical interest only, are the "Exhaustion Theory" of Pasteur and the "Retention Theory" of Nencki,¹³ Chauveau, and others.

Pasteur's views, defended for a time also by Garré, held that the growth of any given variety of bacteria in the animal body exhausted certain specific nutritive substances necessary for this growth. Subsequent lodgment in the same body was impossible owing to the absence of proper nutrient material. It is interesting to note, as Kolle¹⁴ points out, that this theory is in principle very similar to the "Atrepsie" idea of Ehrlich advanced in explanation of species immunity to cancer.

The hypotheses of Chauveau, of Nencki, and others were the converse of those of Pasteur. They were based purely on inference, assuming that conditions occurring in the test tube could be applied also to those existing in the animal body. Baumann¹⁵ had shown that, among other things, phenol was produced as a result of bacterial putrefaction. Nencki had noticed the inhibition of bacteria in culture by the products of their own metabolism. Wernicke,¹⁶ too, had demonstrated the presence of phenol, phenylacetate, skatol, and other aromatic compounds harmful to bacteria in putrefying mixtures. The reasoning which formulated the so-called "Retention Theory," therefore, was the following: Bacteria growing in the animal body produce certain substances peculiar to their own metabolism, which eventually lead to inhibition of their growth. By the retention of these products the animal is rendered immune. Chauveau's adherence to this theory was largely based on the fact that he had observed immunity in the lambs born of Algerian ewes which had recovered from anthrax shortly before or during parturition. He explained this by a transference of the retention products from mother to offspring. As a matter of fact the observation could just as well have been utilized as support for the Exhaustion Theory.

Both the theory of "Exhaustion" as well as that of "Retention" could not long withstand experimental criticism. Theories which were not so easily disproved and which have given rise to much investigation are the "Alkalinity Theory," first formulated by v. Beh-

¹³ Nencki. *Jour. f. prakt. Chem.*, May, 1879, cited from Sirotnin, *Zeitschr. f. Hyg.*, Vol. 4, 1888.

¹⁴ Kolle in "Kolle u. Wassermann Handbuch," 2d Ed., Vol. 1.

¹⁵ Baumann. *Zeitschr. f. physiol. Chem.*, Vol. 1.

¹⁶ Wernicke. *Virch. Archiv*, Vol. 78.

ring,¹⁷ and the "Osmotic Theory" of Baumgarten.¹⁸ In the former an attempt was made to demonstrate a parallelism between blood alkalinity and bactericidal action—the latter was based on the supposition that the destruction of bacteria in the body was largely due to harmful osmotic conditions. Neither of these theories was long seriously maintained. Behring himself took an active part in the subsequent development of our present views. Baumgarten¹⁹ still clings to his own opinion in a modified way, in that he maintains that the only effect produced by specific antibodies upon cells—bacterial or otherwise—is that they change the permeability of the cell membranes and render them more vulnerable to osmotic injury.

However crude or vague these theories may seem to us now, it must not be forgotten that they were conceived at a time when no knowledge had been gained regarding specific "antibodies." The phagocytic powers to which Metchnikoff attributed natural immunity and the bactericidal powers of the blood, regarded in the same light by the Flüge school, were general properties possessed by many animals toward many different micro-organisms. That immunization could specifically increase these functions toward the particular micro-organisms used for treatment seemed indicated by the experiments of Nuttall in which higher bactericidal power was found in the blood of anthrax-immune calves than in that of normal animals. However, no definite and conclusive work on the specific increase of measurable serum or cell properties was available.

This great advance, giving new energy and pointing out new paths of investigation, came in 1890-1892 with the publication of the work of Behring and his collaborators, Kitasato and Wernicke, on immunity to diphtheria and tetanus. As we have indicated in a preceding paragraph, the fundamentally important points of this work were as follows:

1. The establishment of the fact that animals may be actively immunized with products of bacterial metabolism—true toxins or exotoxins.

2. The discovery that such active immunity was dependent upon specific antibodies formed in the treated animal and circulating freely in the blood; and,

3. That, by the transfer of the blood or the blood serum containing these specific antibodies other normal animals could be passively protected—not prophylactically only, but even after active disease had set in.

These observations were rapidly confirmed for tetanus by Tizzoni and Cattani, and by Vaillard, and, similar but less successful attempts at passive immunization were made in other diseases by

¹⁷ v. Behring. *Centralbl. f. klin. Med.*, 1888, No. 38.

¹⁸ Baumgarten. *Berl. klin. Woch.*, 1899, 1900.

¹⁹ Baumgarten. *Lehrbuch, etc.*, 1912.

Foa, Emmerich, Bouchard, and many others. The discovery of passive immunization established the fact of specific alteration of the blood by active immunization, and represented, for the time, a distinct triumph for the humoral hypothesis.

Summarizing the knowledge of immunity as it stood at the close of this period, Behring says: "In the case of natural immunity no generally applicable explanation has as yet been found. (By this he referred to the lack of complete parallelism between natural immunity and either the bactericidal or the phagocytic activities.) For artificial immunization, however, it has now been shown, in a number of carefully studied infections, that we can surely attribute it to properties of the cell-free blood."

Within a very short time after Behring and Kitasato's first paper Ehrlich²⁰ demonstrated that the principle discovered by them was not limited to bacterial poisons. He was investigating immunization against ricin in mice, and showed that here, too, the blood of the immune animals contained a body which would antagonize the toxic action of ricin, and which, injected into normal mice, would passively protect them. He spoke of this blood constituent as "anti-ricin."

It is natural that extensive generalization followed these discoveries. However, while it was found that the blood of all actively immunized animals possessed a certain degree of protective power for normal individuals, it was soon shown that this was not due in all cases to antagonism to the bacterial poisons on the part of the immune blood serum. In immunity to the *Vibrio Metchnikovi*—in pneumococcus and cholera immunity—Sanarelli,²¹ Isaëff,²² Pfeiffer and Wassermann,²³ and a number of others showed that here, unlike diphtheria and tetanus, the protective power of the immune serum did not rest on "antitoxic" properties, but rather on antagonism to the bacteria themselves. It soon became definitely established that antitoxic immunity resulted only in the cases of those bacteria in which a true soluble exotoxin was produced, and where the disease following infection was primarily due to the absorption of these poisons. The antibodies incited in the blood of toxin-immune animals were therefore spoken of by Behring and Ehrlich as "*antitoxins*" and their action—after a number of false hypotheses—was finally recognized as a direct neutralization of the bacterial poisons.

The strict specificity of these antibodies was, from the first, clear to v. Behring, who observed that diphtheria-immune serum and tetanus-immune serum acted each upon its respective toxin only. It was recognized at the same time that the passive immunity produced

²⁰ Ehrlich. *Deutsche med. Woch.*, No. 32, 1891.

²¹ Sanarelli. *Ann. Past.*, Vol. 7, 1893.

²² Isaëff. *Ibid.* and *Zeitschr. f. Hyg.*, Vol. 16, 1894.

²³ Pfeiffer and Wassermann. *Zeitschr. f. Hyg.*, Vol. 14, 1893.

by injecting antitoxic sera is almost immediately established; that, by proportionately increasing the amount of antitoxin, immunity can be produced against any amount of toxin; and that this passive or transferred immunity is of relatively short duration.

The antitoxins, then, as we shall see in the more detailed analysis of their action (in chapter V), are specific poison-neutralizing antibodies formed in the blood of animals immunized with a true bacterial toxin or exotoxin—conferring resistance or immunity, not by influencing the bacteria, but by rendering innocuous the specific bacterial poisons.

The therapeutic successes of passive immunization achieved with tetanus and diphtheria very naturally led to a careful inquiry into the antitoxic properties of the blood of animals immunized with all known pathogenic bacteria and bacterial products, and with many poisons of animal and vegetable origin.

Contrary to earlier expectations, however, the list of bacteria against which antitoxic immunity can be achieved has remained relatively small, limited in fact, as we have previously stated, to those species which produce a soluble exotoxin. The inciting of a specific neutralizing antibody (antitoxin), however, is also a property of many other substances of proteid nature which are for this reason classified biologically with the true toxins or exotoxins. In fact, the one absolutely constant attribute which defines our conception of the “true toxins” and the substances classified with them is their antitoxin-inciting power. We classify a bacterial product as a “toxin” or “exotoxin” only if it incites a neutralizing “antitoxin” in the serum of an immunized animal.

The first discovery of a non-bacterial antitoxin-stimulating substance was, as we have stated, that of ricin by Ehrlich,²⁴ 1891, and this was soon followed by similar determinations for abrin and robin—other vegetable poisons. In 1894 Calmette,²⁵ and Physalix and Bertrand²⁶ extended the principle to poisons of animal origin by demonstrating antitoxin formation against snake poison. And that similar specific neutralizing bodies were formed in response to immunization with ferments was shown in 1900 by Morgenroth.²⁷

The more important individual substances which may be biologically grouped together because of their property of inciting a specific antitoxin (or toxin-neutralizing body) in the blood of immunized animals may be tabulated as follows:

Diphtheria toxin—(*loc. cit.* Behring & Wernicke).

Tetanus toxin—(*loc. cit.* Behring & Kitasato).

²⁴ Ehrlich. *Deutsche med. Woch.*, 1891; *Fortschr. d. Med.*, 1891, 1897.

²⁵ Calmette. *Ann. Past.*, Vol. 8, 1894.

²⁶ Physalix and Bertrand. *Compt. rend. de la soc. de biol.*, 1894.

²⁷ Morgenroth. *Centralbl. f. Bakt.*, 26, 1899.

The Toxin of the Bacillus of Symptomatic Anthrax—(Grassberger & Shattenfroh, *Münch. Med. Woch.*, 1900, 1901 and 10 c. cit.).

The Toxin of the *Bacillus Botulinus*—(Kempner, *Zeitschr. f. Hyg.*, Vol. 26, 1897).

The Toxin of the *Bacillus Pyocyaneus*—(Wassermann, *Zeitschr. f. Hyg.*, Vol. 22, 1896).

The Toxin of the *Dysentery Bacillus* (?) Shiga-Kruse type—(Kraus u. Doerr, *Wien. klin. Woch.*, 1905).

The leukocyte poison of the *Staphylococcus pyogenes aureus*, *Leucocidin*—(Denys & Van de Velde, *La cellule*, 1895).

The Hemolytic Poisons of Various Bacteria (see Pribram in "Kraus und Levaditi Handbuch," Vol. II, p. 223).

Proteolytic Ferments of the Hog Cholera Bacillus (De Schweintz, *Medical News*, 1892).

The Toxin of the *Cholera Spirillum* (?) Brau & Denier, *Compt. rend. de l'acad. des sc.*, 1906, Kraus, *Centralbl. f. Bakt.*, 1906, and *Wien. klin. Woch.*, 1906).

Ricin—(Ehrlich, *loc. cit.*).

Abrin—(Ehrlich, *loc. cit.*).

Krotin—(Ehrlich, *loc. cit.*).

Snake venom—(Calmette, *loc. cit.*).

Spider poison—(Sachs, "Hoffmeister's Beiträge," 1902, and Ehrlich, "Gesammelte Arbeiten," etc.).

Lab. enzyme—(Morgenroth, *loc. cit.*).

Pepsin—(Sachs, *Fortschr. d. Med.*, 1902).

Trypsin—(Achalmé, *Ann. Past.*, 1901).

Leukocytic ferments Leukoprotease—(Jochmann & Müller, *Münch. med. Woch.*, 1906).²⁸

The period of investigation which was initiated by the discovery of the specific antitoxins was replete with efforts to determine true toxins and, consequently, antitoxic immunity for all pathogenic bacteria. We have already mentioned that in many cases these efforts were futile—the bacteria in question being found to secrete no exotoxin and the immunity established against them developing without the formation of demonstrable antitoxin. Metchnikoff²⁹ showed this to be the case with hog cholera as early as 1892, and the investigations of Sanarelli, Isaëff, and Pfeiffer and Wassermann pointed in the same direction.

Perhaps the clearest definition of the conditions prevailing during immunization of animals with non-toxin-forming bacteria was that formulated at this time by Pfeiffer. The importance of the bactericidal power of serum, as discussed before this by Flüge, Nuttall, and others, had dealt largely with variations of this general property in relation to natural immunity, but had failed to recognize clearly a specific increase in these powers during active immuniza-

²⁸ This list includes all the important antitoxin-ineiting substances. For a more complete tabulation see Wassermann in "Kolle u. Wassermann Handbuch, etc.," Vol. IV, 1st Ed., p. 498. Our own list is adapted from the one there given.

²⁹ Metchnikoff. *Ann. Past.*, 1892.

tion. Pfeiffer with Wassermann³⁰ had studied the pathogenicity of cholera spirilla for guinea pigs, and had come to the conclusion that the animals died of toxemia (and not of bacteriemia, as claimed by Gruber and Wiener), and that this toxemia was due to the liberation of poisons from the dead bodies of cholera vibrios, killed by the serum of the infected animals. Pfeiffer³¹ now showed that the injection of cholera spirilla killed with chloroform brought about a toxemia identical with that following inoculation with living cultures. He further determined that the resistance of animals against cholera was due to the bactericidal effects of the serum, which killed the injected cholera spirilla, and not to any poison-neutralizing property.

Isaëff,³² one of Pfeiffer's pupils, continuing this work, expresses his own and Pfeiffer's conceptions as follows: "Guinea pigs vaccinated against cholera, in spite of high immunity to infection with living spirilla, do not develop any immunity to cholera [endo]³³ toxins. The blood of immunized guinea pigs possesses no antitoxic properties. The maximal dose of cholera 'toxin' which immunized guinea pigs can withstand is not higher than that which can be borne by normal animals, and but slightly higher than the maximal dose of living spirilla, which they can survive. The blood of cholera-vaccinated guinea pigs possesses strong specific protective powers. The same specific immunizing properties are demonstrable in the blood of cholera convalescents toward the end of the third week of the disease."

The path was thus cleared for a definite conception of cholera immunity, and this was formulated, in their next communication, by Pfeiffer and Isaëff.^{34 35 36} In this paper they showed that the cholera spirilla injected into the peritoneum of a cholera-immune guinea pig were subjected to a rapid dissolution, a process which could be observed by taking small quantities of exudate out of the peritoneum, at varying intervals, with capillary pipettes. No such dissolution occurred in normal pigs or with normal serum. But the same rapid swelling, granulation, and, finally, dissolution occurred when the spirilla were injected into the peritoneal cavity of a normal guinea pig, together with the serum of an immunized animal. The process took place apparently without the coöperation of the leukocytes or other cells, and was absolutely specific. For instance, no "lysis" occurred when the vibrios "Nordhafen," "Massauah," and other cholera-like organisms were injected into cholera-immune pigs,

³⁰ Pfeiffer and Wassermann. *Zeitschr. f. Hyg.*, Vol. 14, 1893; also Pfeiffer, *Zeitschr. f. Hyg.*, Vol. 16, 1894.

³¹ Gruber and Wiener. *Archiv f. Hyg.*, Vol. 15, 1893.

³² Isaëff. *Zeitschr. f. Hyg.*, Vol. 16, 1894.

³³ Bracketed word our own.

³⁴ Pfeiffer and Isaëff. *Zeitschr. f. Hyg.*, Vol. 17, 1894.

³⁵ Pfeiffer. *Ibid.*, Vol. 18, 1894.

³⁶ Pfeiffer and Isaëff. *Deutsche med. Woch.*, No. 13, 1894.

but took place regularly when true cholera strains, from various sources, were used in the experiment. The immunity of cholera-treated animals, therefore, was found to be an antibacterial and not an antitoxic one. Cholera spirilla introduced into a normal animal were permitted to multiply and accumulate until a sufficient number were present to furnish, upon cell death, a fatal dose of poison. In immunized animals the small quantities of bacteria first introduced succumbed rapidly to the lytic properties of the serum and accumulation was prevented.

By these experiments, now commonly spoken of as the "Pfeiffer Phenomenon," it was definitely proved that active immunization with bacteria incites in the serum of the treated animal a potent increase of bactericidal properties—an increase which is entirely specific in that the bactericidal power toward bacteria other than those employed in the immunization does not exceed the normal. The immunity in these cases, then, is not *antitoxic*, but rather "*antibacterial*," and depends on the development, in the immune sera, of antibodies quite distinct from the "*antitoxins*." These immune serum constituents were spoken of by Pfeiffer as "*bacteriolysins*" or "specific bactericidal substances."

Not long after the discovery of the specific bacteriolysins another property of immune sera was described by Gruber and Durham.³⁷ They had been studying bacteriolytic phenomena with colon and cholera organisms, and noticed that these bacteria were rapidly agglomerated and gathered in small clumps when emulsified in homologous immune serum. Similar clumping had indeed been described before. Metchnikoff, Isaëff, Washburn, and Charrin and Roger had described it on various occasions, but had not recognized it as a specific property of immune serum.³⁸ Gruber and Durham studied it carefully, determined that it was present to a degree roughly proportionate to the degree of immunization attained, and that its specificity was such that it could be utilized for bacterial differentiation. They believed that the substances in the immune serum responsible for this agglutination were independent of other serum constituents and applied to them the term "*agglutinins*."

The problems of immunization had now considerably expanded and the nature of the new serum reactions was assiduously studied. Primarily the phenomenon of agglutination was regarded as a part of the struggle of the body against the living bacteria and Gruber himself believed that it depended upon a swelling or "*klebrig werden*" of the micro-organisms which tended to cause their sticking together, and rendered them more readily amenable to the action of the bactericidal powers of the serum. Bordet,³⁹ however, early con-

³⁷ Gruber and Durham. *Münch. med. Woch.*, 1896.

³⁸ For references see chapter on Agglutinins.

³⁹ Bordet. *Ann. Past.*, 1896.

ceived the process as a physical phenomenon in which the bacteria themselves were entirely passive, and, indeed, Widal⁴⁰ soon demonstrated that bacteria killed by heat were equally as agglutinable as the living germs.

This naturally suggested that the reaction between specific agglutinating serum and bacteria was based on individual peculiarities of the bacterial proteins, and it occurred to Kraus,⁴¹ accordingly, to investigate whether or not the immune sera would cause any sort of reaction when mixed with the dissolved body substances of homologous bacteria. Working at first with cholera and plague, he prepared solutions of bacterial proteins, both by allowing broth cultures to stand for varying periods and by emulsifying agar cultures in alkaline broth. The extracts were then filtered through Pukal filters to remove the bacterial bodies. When the sera of immunized animals were added to these clear filtrates—cholera serum to cholera filtrate, and plague serum to plague filtrate, slight turbidity developed and was followed within twenty-four hours by the formation of small flakes. In other words, it was found that the mixture of a clear filtrate of a bacterial culture with the serum of an animal immunized against these bacteria resulted in the formation of a precipitate. The reaction was found to be as strictly specific as that of agglutination.

Although, from the beginning, Paltauf⁴² attempted to associate the phenomena of agglutination and precipitation, the property of precipitating homologous culture filtrates was attributed by Kraus and others to specific antibodies in the immune sera, distinct and independent of those previously described, and spoke of them as "*precipitins*."

The discovery of the various "antibodies" so far discussed resulted from the study of the direct action of blood serum upon bacteria and bacterial products. This did not, however, completely deflect the attention of investigators from the unquestionable importance of phagocytosis in the defence of animals against bacterial invasion. Metchnikoff and his school continued diligently to pursue this other phase of the study of immunity and, although the increasing knowledge of serum antibodies continued to strengthen the premises of the purely humoral point of view, it had still to be admitted that in some diseases—particularly anthrax and the pyogenic coccus infections, phagocytosis must largely be held responsible for recovery. It was found, moreover, by the later investigations of Denys, Wright, Neufeld, and others that phagocytosis in immunized animals was far more extensive and efficient than in normal ones, and that

⁴⁰ Widal. *La semaine médicale*, No. 5, 1897.

⁴¹ Kraus. *Wien. klin. Woch.*, No. 32, 1897.

⁴² Paltauf. "Discussion of Kraus' Paper," *Wien. kl. Woch.*, No. 18, 1897. p. 431.

this depended on specific constituents of the immune serum which rendered the bacteria more amenable to the phagocytic action of the cells. These further antibodies we will discuss in a subsequent chapter, under the terms "*opsonins*" and "*bacteriotropins*," designations applied to them by their discoverers.

We have thus reviewed briefly the various specific properties which develop in the serum of an animal when it is systematically treated (actively immunized) with bacteria or bacterial products. These serum activities have been attributed to the development in the serum of substances which we speak of as "*antibodies*."

In our discussion of the first of these antibodies, antitoxin, we call attention to the fact that the principle discovered in the case of bacterial toxins was rapidly extended to vegetable poisons, snake venom, spider poison and enzymes. It was found that the power of inciting antitoxins when injected into animals was an attribute belonging to a large group of substances in nature, and not limited to bacteria alone. A similar generalization of conception has been possible with other antibodies. Specific lysins, agglutinins, and precipitins may be produced by the treatment of animals with many substances not of bacterial nature.

The first observation of this kind was made almost simultaneously by Bordet⁴³ and by Belfanti and Carbone.⁴⁴ They observed that the serum of an animal that had been treated with the red cells of another species acquired the power of laking these cells. That the normal serum of one species is often toxic to, and causes the laking of, the erythrocytes of another species is an observation that dates back to the earliest experiments on transfusion, and had been studied in considerable detail by Landois as early as 1875. The phenomenon possesses much interest in its bearing on the problems of anaphylaxis and will be discussed more particularly in that connection. We mention it in this place to show that, like bactericidal bodies, "hemolytic" (erythrocyte laking) properties may be present in normal sera, though irregularly and by no means occurring in every species of animal. Incidentally it may be stated that this is true also of agglutinins and of opsonins which may be found in considerable amounts in normal sera. Of precipitins, however, this does not seem to be true.

By the work of Bordet it was found that "hemolysins" could be specifically⁴⁵ incited in an animal by systematically treating it with

⁴³ Bordet. *Ann. Past.*, Vol. 12, 1898.

⁴⁴ Belfanti and Carbone. *Giorn. della R. Acad. di Torino*, July, 1898.

⁴⁵ By the use of the word specific in this case we imply that an animal immunized with any given variety of red blood cells will form hemolysins for this variety only. Thus an animal treated with ox blood will form ox blood hemolysins only, and his serum, though strongly hemolytic for ox blood, will not lake sheep cells, dog cells, human cells, etc.

the red blood cells of another species. Apart from the great interest attaching to this discovery in itself, it has had a very profound influence upon investigations on immunity generally, since it has furnished a method of studying lysis far more simple and easily controlled than is the analogous phenomenon of bacteriolysis. And since, in fundamental principles, bacteriolysis and hemolysis are essentially alike, much of our knowledge regarding the former has been arrived at by experiments upon the latter. The specific *hemolysins*, then, are antibodies formed in response to "immunization" with red blood cells, analogous to the similarly produced "bacteriolysins." Because both of these antibodies exert definite injury upon cells, we speak of them by the group names of "*cytolysins*" or "*cytotoxic*" substances.

The discovery of hemolysins naturally suggested the use of other cells, and the following years brought forth many reports of further specific cytotoxins. In 1899, Metchnikoff,⁴⁶ and very soon afterward Landsteiner,⁴⁷ described specific "*spermatotoxins*" which appeared in the blood of animals treated with spermatozoa. Von Dungern⁴⁸ obtained analogous substances by injecting ciliated epithelium from the trachea. Neisser and Wechsberg⁴⁹ produced "*leukotoxin*" by injecting leukocytes; Delezenne⁵⁰ produced "*neurotoxin*" and "*hepatotoxin*," and Surmont,⁵¹ *pancreas cytotoxin*. Subsequent years have added to these "*gastro-toxin*" (Bolton),⁵² *thymotoxin* (Slatineau),⁵³ *adrenal cytotoxin* (Gildersleeve),⁵⁴ *placental cytotoxin* (Frank),⁵⁵ *corpus luteum cytotoxin* (Miller),⁵⁶ and a number of others. In fact, as Roessle⁵⁷ puts it, in a review of the literature, there is no organ in the body for which it has not been claimed that specific cytotoxins can be formed by the injection of homologous macerated tissues.

Recent critical study of these organ-cytotoxins has revealed, however, that the specificity of a serum produced with the tissues of one organ is not strictly limited to this organ alone, and that the serum may injure other organs as well. It is true, indeed, that there are certain cells and tissues in the body such as the spermatozoa, the tissues of the testicles, the ovary, the lens of the eye, and, possibly,

⁴⁶ Metchnikoff. *Ann. Past.*, Vol. 13, 1899.

⁴⁷ Landsteiner. *Centralbl. f. Bakt.*, Vol. 25, p. 549, 1899.

⁴⁸ Von Dungern. *Münch. med. Woch.*, p. 1228, 1899.

⁴⁹ Neisser and Wechsberg. *Zeitschr. f. Hyg.*, Vol. 36, 1901.

⁵⁰ Delezenne. *Ann. Past.*, 1900; *Compt. rend. de l'acad. des sc.*, 1900.

⁵¹ Surmont. *Compt. rend. de la soc. de biol.*, 1901.

⁵² Bolton. *Lancet*, 1908.

⁵³ Slatineau. Cited from Roessle, *loc. cit.*

⁵⁴ Gildersleeve. Cited after Roessle.

⁵⁵ Frank. *Jour. Exp. Med.*, 1907.

⁵⁶ Miller. *Centralbl. f. Bakt.*, 47, 1908.

⁵⁷ Roessle. "Lubarsch und Ostertag," Vol. 13, 1909.

the placenta which have chemical characteristics so well defined and individual that the cytotoxic sera induced by them have definite organ specificity. The same to a more limited extent seems true of kidney substance (Pearce). In most cases, however, in which originally a specific cytotoxin was claimed, it has been possible to show subsequently that the apparently selective injury was due not to organ specificity alone but to the fact that the injection of tissue-macerates, even when sufficiently freed from blood, induced the formation of considerable amounts of hemagglutinins and hemolysins.

Pearce⁵⁸ expresses it as follows: “. . . it is evident that the cells or the various organs of the body, while differing in morphology and function, have certain (receptor) characteristics in common, and that one type of cell may therefore produce antibodies affecting several cells of differing morphology, but with like (receptor) groups. This is shown by the sera prepared from washed liver, kidney, pancreas, and adrenal, all of which may agglutinate and hemolyze red blood cells and may cause degenerative changes also in the liver and the kidneys. Some of these cytotoxic sera have no effect upon organs for which they are supposed to have a morphological affinity, but exert a powerful lytic influence upon other cells. Aside from nephrotoxin, which has a distinct injurious action upon renal epithelium, the various cytotoxins studied (kidney, liver, pancreas, and adrenal) have no specific action in the morphological sense.”

This opinion seems to be in harmony with that of most observers who have studied the problem recently, at least as regards most of the organ cytotoxins. Much of the promised light upon pathological processes—looked for when cytotoxins were first studied, has faded, moreover, since it has been found that cytotoxins cannot be produced by injection into an animal of cells, tissues, or fluids from its own body. “Autocytotoxins” in general cannot be produced, a question discussed at greater length in the chapter on lysis, in connection with Ehrlich’s work on the isolysins.

The work outlined in the preceding paragraphs had thus extended the principles of antitoxin and lysin production beyond the scope of pure bacteriology, and had shown them to possess the significance of general biological laws. Similar generalization was soon attained in the case of the agglutinins and in that of the precipitins. In the former, the nature of the reaction limited it to observations upon cells in suspension, and, in connection with the earlier experiments upon hemolysis it was soon discovered that the erythrocytes were often clumped before lysis could take place, when brought together with a hemolytic serum of moderate or feeble potency, or when solution, for other reasons, was delayed.

⁵⁸ Pearce. *Jour. of Med. Res.*, N. S., Vol. 7, 1914, p. 13.

The first observations on the general significance of the precipitin reaction we owe to Tschistovitch⁵⁹ and to Bordet.⁶⁰ Tschistovitch was studying the toxic action of eel serum upon rabbits. This serum, as Kossel⁶¹ had shown, is toxic for rabbits and possesses the property of causing hemolysis of rabbit erythrocytes. Its similarity to ricin, in this respect, stimulated attempts to produce an antitoxic substance against eel serum, even as Ehrlich had produced an antiricin. In the course of such experiments Tschistovitch observed that, when eel serum was mixed with the serum of a rabbit which had received several injections of this substance, the mixture became rapidly opalescent and soon a flocculent precipitate was formed. Coincident with this discovery Bordet made a similar observation. He had injected chicken blood into rabbits in the course of experiments upon hemagglutination. He found that the serum of the rabbits so treated acquired the property not only of producing hemolysis and hemagglutination of chicken cells, but also of giving a precipitate if mixed with chicken serum.⁶² Soon after this precipitins were produced by injecting rabbits with milk (Bordet), egg albumen (Ehrlich, Uhlenhuth), and many other substances, and the specificity of such reactions was demonstrated by Fish,⁶³ Wassermann and Schütze,⁶⁴ Uhlenhuth, and many others.

It is apparent from the preceding paragraphs that the discovery of specific antitoxins merely constituted the first step in the formulation of a fundamentally important biological law. There is, then, a large group of substances of animal and vegetable origin which call forth the formation of specific reacting bodies when injected into animals. In order to elicit this response it is necessary that these substances shall penetrate to the physiological interior of the body in a relatively unchanged condition. For this reason any form of injection, subcutaneous, intravenous, or into a serous cavity, is followed, with regularity, by antibody formation, whereas feeding or other means of intrainestinal administration is negative in result, unless abnormal conditions prevail which permit entrance into the blood before the digestive enzymes have decomposed the ingested materials.

The substances with which antibody-formation may be induced are collectively spoken of as "*antigens*." Within this group we may distinguish two main subdivisions, indicated in our preliminary dis-

⁵⁹ Tschistovitch. Cited by Bordet, *loc. cit.*, and also *Ann. Past.*, 13, 1899.

⁶⁰ Bordet. *Ann. Past.*, Vol. 13, 1899.

⁶¹ Kossel. *Berl. klin. Woch.*, No. 7, 1898.

⁶² This, we know now, was due to the fact that the blood cells injected were not washed free of chicken serum. Thus chicken serum precipitin was formed as well as were hemagglutinin and hemolysin.

⁶³ Fish. *St. Louis Med. Jour.*, 1900. Cited from Uhlenhuth.

⁶⁴ Wassermann and Schütze. *Deutsche med. Woch.*, No. 30, 1900. Vereinsbeilage.

cussions. The first of these is composed of the true bacterial toxins—the vegetable poisons ricin, abrin, krotin and robin, snake venom, the enzymes, and other substances grouped with these on page 87. These antigens have certain characteristics which render them comparable to ferments, and they induce in the animal body the formation of specific neutralizing antibodies—(antitoxin, antivenin, anti-enzyme)—which inactivate their respective antigens when mixed with them in proportionate quantities, either in the test tube or in the animal body. This characteristic alone separates them sharply from other antigens. The remaining antigenic materials do not induce antitoxin-like neutralizing substances, but call forth specific lytic, agglutinating, precipitating, or opsonic properties.

Since the phenomenon of antibody formation is not at all limited to bacteria or bacterial derivatives, it cannot be looked upon merely as a mechanism existing for the primary purpose of protecting the body against infectious disease. This latter function is important, indeed, but is probably incidental to the broader significance of the processes.

In the course of normal existence substances which are not directly assimilable as such—foreign proteins, for instance—do not penetrate directly into the blood and tissues. Taken into the alimentary canal, they are first hydrolized into peptons, albumoses, polypeptids, and probably amino-acids before absorption, to be reconstructed from these cleavage products (“Bausteine” is Abderhalden’s expression for the amino-acids) into protein biologically identical with that of the tissues. Digestive and other accidents, however, on numerous occasions during life permit the direct entrance of these materials unchanged or insufficiently changed into the circulation. It is probably by the action of digestive powers of the serum—or, in the case of the entrance of undissolved foreign particles, by the activity of the phagocytic cells—that such substances are then disposed of and assimilated. For each particular variety of substance (antigen) a specific mechanism is called into play, and when this mechanism is repeatedly called upon—as in successive injections of foreign proteins—this mechanism, whatever it may consist of, is enhanced in efficiency—i. e., increased in quantity. How this increase of specific antibodies is theoretically conceived we will discuss later in connection with Ehrlich’s side-chain theory.

The phenomena of antibody formation against bacteria on this basis may be taken to constitute, then, a mechanism for the digestion and disposal of a foreign protein which has penetrated into the tissues and, because of its living state, increases within the body by multiplication, furnishing progressive stimulation to the antibody-producing function. Infectious disease, therefore, from this point of view may be looked upon as an invasion of the body by a living foreign protein which must be assimilated and disposed of; which,

in some cases, has a primary toxicity per se; and which is variously distributed among the organs and tissues according to the biological peculiarities of the particular micro-organism in question. This general conception will become more clear as we analyze the phenomena associated with the individual antibodies. It is, of course, quite plausible as far as it refers to the phagocytic functions, or even bacteriolytic and cytolytic phenomena. It has been less clear in connection with the agglutinins and precipitins in which a direct defensive or bacteria-destroying value is not apparent. However, in our discussions of these phenomena we will have occasion to point out many reasons for assuming that, even in these phenomena, there are features which fall into direct correlation with the views we have just expressed.

The substances which possess antigenic properties—that is, which give rise to antibody production—with the exception of a few isolated and contested cases, are all of them protein in nature. Well-trained chemists have exerted themselves to purify antigenic substances, in attempts to determine the particular fractions of the complex protein molecule upon which the antigenic properties depend. In the course of such work a number of men claim to have obtained a truly antigenic substance which no longer gave protein reactions. The instance most frequently cited is Jacoby's⁶⁵ announcement of a protein-free ricin. Jacoby worked with an apparently very impure "Ausgangsmaterial" consisting of commercial ricin, which he digested for five weeks in trypsin solution. At the end of this time he obtained a ricin which still possessed the properties of the original castor-bean extract, but no longer gave protein reactions. His "purified ricin," however, was quickly destroyed by further trypsin digestion, and more recent work by Osborne, Mendel, and Harris⁶⁶ appears to have fully refuted Jacoby's results. They found the purified ricin identical with the coagulable albumin of the castor bean, and found that tryptic digestion destroys the characteristic ricin properties.

Less easily refuted have been the careful experiments of Ford⁶⁷ upon the active principle of a mushroom (*Amanita phalloides*) and upon that of the poison-ivy plant—(*Rhus toxicodendron*). These substances, he claims, are non-protein. In the case of *Amanita phalloides* Abel and Ford⁶⁸ have shown it to be a glucosid, and similar structure has been claimed for Rhus by Syme.⁶⁹ Yet with both of these substances Ford has succeeded in producing specific

⁶⁵ Jacoby. *Arch. f. exp. Path. u. Pharm.*, Vol. 46, 1901.

⁶⁶ Osborne, Mendel, and Harris. *Am. Jour. of Physiol.*, 1905, Vol. 14.

⁶⁷ Ford. *Jour. of Inf. Dis.*, Vol. 3, 1906; Vol. 4, 1907.

⁶⁸ Abel and Ford. *Jour. Biol. Chem.*, 1907.

⁶⁹ Syme. *Johns Hopkins Thesis*, 1906.

antitoxins. Rabe ⁷⁰ has recently questioned the results of Abel and Ford with *Amanita phalloides*. He believes that the poison with which Ford worked is not a glucosid, but is of protein nature. In the case of Rhus, however, Ford's conclusions have not, to our knowledge, been challenged.

With these and a few other less important exceptions, however, observers have uniformly concluded that antigenic property and protein structure are inseparably associated. All procedures by which proteins have been hydrolized into their simpler fractions, chemical splitting, tryptic or peptic digestion have in every case resulted in a simultaneous loss of protein reaction and antigenic property.

Many attempts have also been made to show a relation between antigenic properties and the lipoid constituents of cells. These endeavors were obviously stimulated by the observation that many lipoids are capable of binding antibodies *in vitro*, and that, in nervous tissues, toxin fixation was in some way related to the richness in lipoids of these structures. Bang and Forsmann ⁷¹ accordingly treated animals with ether extracts of red blood cells—claiming that this resulted in the production of hemolysins. And these results have been confirmed by Landsteiner and Dautwitz.⁷² The latter, however, suggest that the hemolysin production may have been induced, not by the lipoidal substances in solution, but by other antigenic substances which had gone into colloidal suspension in the ether extracts. Much similar research on the antigenic nature of lipoids has been done, but, after reviewing this very thoroughly, Landsteiner comes to the conclusion that no definite proof of the antigenic nature of any pure lipoid has so far been presented. The problem is experimentally complicated by the fact that, as Landsteiner ⁷³ suggests, the antigen may often be present as a lipoid-protein combination, and as such go into solution or fine emulsion in the organic solvents; also the lipoids possess the curious property of altering the solubilities of proteins and other substances by their presence.

Summarizing our present knowledge of the chemical nature of antigens, then, we must conclude that, with the exception of Ford's glucosids, no protein-free antigens have been thus far demonstrated.

In the light of this fact it is all the more remarkable that antigen-antibody reactions are specific. For we possess no chemical methods by which one variety of protein can be distinguished from another. And yet the serum antibodies produced with each species of bacteria

⁷⁰ Rabe. *Zeitschr. f. exp. Path. u. Therap.*, Vol. 9, 1911.

⁷¹ Bang and Forsmann. *Hofm. Beitr.*, 1906; *Centralbl. f. Bakt.*, 40, 1906.

⁷² Landsteiner and Dautwitz. *Hofm. Beitr.*, 9, 1907.

⁷³ Landsteiner. "Wirken Lipoide als Antigene?" *Weichardt's Jahresbericht*, Vol. 6, 1910.

react with this species only—and the hemolysins, agglutinins, or precipitins produced by the injection of bacterial, cellular, or serum proteins react respectively only with the particular variety employed in their production. This indicates that each of these antigens—of almost unlimited number—must possess a chemical structure individually characteristic and different from all the others. It is by means of the biological reactions, indeed, that we can detect protein in dilutions far beyond the reaction-sensitiveness of chemical tests and can distinguish between varieties of protein when the chemical methods will indicate only protein in general. Our knowledge of the chemical constitution of protein has not yet advanced to a point at which specificity can be based upon definite variations of chemical structure, and the complexity of the problem is such that it does not seem likely that we can hope in the near future to attain such knowledge. We can merely accept it as a fact that the antibody produced with one protein differs materially from that produced with another, and that this is a definite indication that the antigen in one case must be chemically different from that in another.

The range of such variations is apparently enormous. For each variety of bacteria or plant, each species of animal, and to a certain extent each individual of the species, possesses certain special antigenic characteristics peculiar to itself. In general there is an underlying antigenic similarity which is peculiar to the species. This is true of bacteria and, in the case of animal and vegetable proteins, an antibody produced with material from an individual of a certain species will react with the protein derived from this species in general. However, that there are also antigenic differences between individuals within the same species is indicated by Ehrlich's experiments on the antibodies produced by injecting the blood cells of one goat into another. And we have further indicated that within the same animal different organs may possess individual antigenic characteristics. Added to this we know that certain special organs like the testicle, the lens, and some others contain antigens which are peculiar to this variety of organ, irrespective of species—a condition spoken of as "*organ specificity*." Thus an antibody produced by injections of the testicular substance of one animal will react with testicular protein from many different species—the specificity here depending upon the organ and not upon the zoölogical relationship.

It is clear, therefore, that there are more different varieties of protein, biologically distinguishable, than there are species of living beings in nature. As Abderhalden⁷⁴ has recently pointed out, this is a conception which it is a little difficult to grasp chemically, since in breaking up different proteins into their "building stones" (*Bausteine*) we encounter again and again the same 20 amino-acids. By a simple arithmetical consideration, however, he shows that merely

⁷⁴ Abderhalden. *Münch. med. Woch.*, No. 43, 1913.

by combining these twenty amino-acids in different groupings an enormous number of isomeric but varying compounds can be formed—even without assuming the additional possibility of quantitative variations. He reasons that 3 “Bausteine”—A, B, and C—could form 6 different structures, A B C, A C B, B C A, B A C, C A B, C B A. Similarly 4 could form 26, and finally 20 could form 2, 432, 902, 008, 176, 640, 000 different compounds.⁷⁵

The analogy between the active immunization of animals with the various antigens and certain chemically well-defined poisons, alkaloids, etc., is so obvious that it has led to much speculation as to a possible similarity in the physiological mechanisms of the two phenomena. As a matter of fact the acquired tolerance for such substances as morphin, atropin, and other alkaloids is not really analogous to the physiological reactions which follow the treatment of animals with bacterial and other proteins, for whatever toxic properties there are in the latter are, as we shall see later, rather the results of the interaction of these injected substances and the reaction products supplied by the cells and fluids of the body. It is at least probable in the light of our modern conceptions that such protein antigens are not toxic per se, in the native state. This, however, will receive detailed consideration in succeeding sections. The analogy of drug tolerance, however, to the acquired immunity against true bacterial toxins and vegetable poisons like ricin, croton, and others is a striking one, since in both classes of poisons there is a gradually developed tolerance for substances toxic in the native state and often very similar in physiological effects (strychnin and tetanus toxin, etc.). In the case of the toxins, however, there is a development of immunity by actual neutralization of the poisonous principle brought about by a specific antibody, which circulates in the blood of immunized animals and man—the process following, within certain limits, the law of multiple proportions. In the case of morphin and other alkaloids no such neutralizing antibodies have as yet been demonstrated.⁷⁶ Whereas toxin immunity is passively transferable from one animal to another with the blood serum, and, *in vitro*, the mixture of the toxin with the immune serum brings about a neutralization of the poison, no such phenomena have been observed, as a general rule, in the case of the alkaloids. We say “as a general rule” since an exception is recorded in the observations of Fleischmann,⁷⁷ who claims to have found antagonistic action to atropin in the blood of normal rabbits, this power being absent from the blood

⁷⁵ We have not repeated the arithmetical labor and take Abderhalden's word for it.

⁷⁶ Hans Meyer and Gottlieb. “Exp. Pharm.” 2d Ed., Neban & Schwartzberg, Berlin, 1911, p. 517.

⁷⁷ Fleischmann. *Archiv f. exp. Path. u. Pharm.*, 62, 1910, cited from Meyer and Gottlieb, *loc. cit.*

of rabbits that had thyroid hypertrophies and were, in consequence, atropin-susceptible. Other observations of a similar significance have been made by Physalix and Contejean⁷⁸ on curare, but have not been confirmed, and the investigations of all other workers on this subject have had negative results. It seems from available evidence that tolerance (immunity) against drugs is due to cellular rather than to serum antagonism.

THE ORIGIN OF ANTIBODIES

The tissue cell, as the ultimate functional unit, must, of course, be looked upon as the source from which originate the various protective constituents of normal and immune sera; and, though perhaps unrecognizable by the coarse tests of morphological investigations, it is in the cells that changes must take place primarily when the animal body is subjected to any one of the processes spoken of as immunization. The exact location of the antibody-forming cells and tissues, in spite of much investigation, is not at all clear, though many data seem to point to the lymphatic organs, the spleen, and the bone marrow as particularly concerned with this process.

Thus Pfeiffer and Marx⁷⁹ exsanguinated animals five days after injections of dead cholera spirilla and found that at this time bacteriolytic antibodies were more concentrated in the spleen than in the blood serum itself. Wassermann's⁸⁰ analogous experiments with typhoid bacilli seemed to show a higher antibody content in spleen, bone marrow, thymus, and lymph nodes than was present in the blood at an early period of immunization. Although these investigations, as well as many others of Castellani,⁸¹ seem, therefore, to indicate a particular association of the special lymphatic organs with antibody formation,⁸² extirpation of the spleen⁸³ before immunization has not prevented animals from responding to injections of bacteria and red blood cells with sharp antibody production. The experiments of Deutsch,⁸⁴ in which reduction of antibody formation resulted in animals in which splenectomy was practiced three or four days after immunization was begun, can hardly be accepted as a conclusion, in the writer's opinion at least, since any severe operation or interference with the normal functions of an animal during the severe physiological strain of active immunization would naturally lead to a less perfect response. That the resistance of animals and man to infection with bacteria is not noticeably diminished by sple-

⁷⁸ Physalix and Contejean. Cited from Meyer and Gottlieb.

⁷⁹ Pfeiffer and Marx. *Zeitschr. f. Hyg.*, Vol. 27, 1898.

⁸⁰ Wassermann. *Berl. klin. Woch.*, p. 209, 1898.

⁸¹ Castellani. *Zeitsch. f. Hyg.*, Vol. 37, 1901.

⁸² Pfeiffer and Marx. *Loc. cit.*

⁸³ I. Levin. *Jour. Med. Res.*, Vol. 8, 1902.

⁸⁴ Deutsch. *Ann. de l'Inst. Past.*, Vol. 13, 1899.

nectomy, moreover, has been variously shown. In unpublished experiments by the writer splenectomized guinea pigs showed no difference from normal animals in regard to their susceptibility to tuberculosis. And though these and similar experiments of other workers with various bacteria are not entirely devoid of interest, their negative results as a matter of fact have no great significance, since our knowledge concerning the true function of the spleen is very incomplete, and it is not impossible that on removal of this organ other elements of the lymphatic system may take over its function in part or as a whole.

Removal of the spleen has not been an extremely unusual procedure in surgery, and there is no evidence to show that patients so treated have been abnormally susceptible to infection thereafter.

Yet, as we have seen, there seems to be an early concentration of antibodies in the lymphatic organs in the course of immunization, and it may well be that an association between the process and these tissues exists which cannot be experimentally demonstrated with absolute certainty.

It is no less likely, however, that similar functions are exerted by the cells of other organs. In fact, it is more than probable that antibodies may be formed anywhere in the body—and that the locality of their production is largely dependent upon the locality in which the antigen is concentrated. Wassermann and Citron⁸⁵ demonstrated this by injecting typhoid bacilli into rabbits intraperitoneally, intravenously, and intrapleurally, and nine days afterward determining the comparative bactericidal strength of blood serum and of aleuronat exudates of pleura and peritoneum in each of the three animals. Their results showed that the bactericidal titre of the intravenously inoculated animal was highest in the blood serum, while that of the intraperitoneally and intrapleurally inoculated animals was highest in peritoneal and pleural exudates respectively. Such experiments point to the possibility of a "local" immunity, that is, a production of antibodies directly by the cells with which the antigen comes into contact in the most concentrated and direct manner. And, indeed, another isolated experiment of the same authors, alone successful of a series of similar attempts, would point in the same direction. Typhoid bacilli were injected subcutaneously into the ear of a rabbit and the ear immediately ligated at its base and kept so for several hours. After nine days the bactericidal titre of the blood serum was determined and the ear amputated. An immediate and rapid drop of antibody contents occurred after the amputation—indicating that the chief source of antibody function had been removed. More striking examples of the same thing are to be seen in the experiments of Römer,⁸⁶ who instilled abrin into a rabbit's eye and found that the

⁸⁵ Wassermann and Citron. *Zeitschr. f. Hyg.*, Vol. 50, 1905.

⁸⁶ Römer. *Arch. f. Ophthal.*, 52, 1901.

retina of the eye developed an antitoxic power against abrin which protected mice against many times the fatal dose, while that of the other eye remained practically inactive.

From these facts, as well as from other observations, it is at least reasonable to believe that antibody formation is by no means a function of special organs and that many cells throughout the body may take part in the process. It is of especial importance to consider this in connection with the possible effects of the treatment of infections by means of bacterial vaccines. If the focus of the infection can possibly become also a local source of antibody production then such treatment may well seem rationally founded, even in generalized acute infections in which no logical basis for such treatment would exist, were the production of antibodies a task for specialized organs like spleen and bone marrow only. The therapeutic phases of this problem are more extensively considered in a later chapter.

It is in this fact also that we must seek the explanation of the apparent local immunity which occurs in certain infections of the skin. Thus it frequently happens that successive crops of boils may afflict different parts of a patient's skin—new ones arising as old ones heal, showing that the process of the limitation and healing of the infected foci is not due to any increase of generalized resistance, but rather to local causes. In the same way, in erysipelas, the process extends along the edges while the original central area of infection is returning to the normal state, and it rarely occurs in adults that the erysipelatous process extends back into the originally infected area.⁸⁷ From these localized laboratories of antibody formation, of course, distribution to the circulation probably takes place and the complete cure of the patient must await a sufficient concentration of these in the body as a whole before further local foci cease to arise.

That the fixed tissue cells of any part of the body can and do take an active part in the local reaction against the invasion of bacteria and other foreign materials is histologically evident. When a more or less insoluble foreign body—a thread of lint, paraffin, agar-agar, or other material—is deposited in the subcutaneous tissues anywhere in the body, and is accompanied by acute infection with bacteria, there is a characteristic tissue reaction which results in the surrounding of the foreign particle by multinucleated cells spoken of as giant cells. In the case of foreign bodies such as those mentioned the process is purely one of local ingestion of the particle which later, if the material remains absolutely insoluble, results in encapsulation by connective tissue. If soluble, however, there may be an eventual digestion of the foreign material by the cell with a subsequent degeneration or splitting up of the giant cell and a return to normal. This also occurs in the case of such infections as those

⁸⁷ In children erysipelas not infrequently returns within a few days over a recently healed area.

due to yeasts or blastomyces, in which, as the writer has seen, the apparent lack of liberation of toxic products gives rise to a purely local giant-cell reaction, adjacent tissue cells remaining undegenerated and apparently unaffected. In the case of infection with bacteria like the bacillus of tuberculosis, the leprosy bacillus, that of rhinoscleroma, and a few others the purely local picture of giant-cell phagocytosis is complicated by secondary reactions arising probably from the liberation of toxic products from the living or dead invaders which both stimulate specific cell reactions and call forth cell degeneration in adjacent tissues, frequently giving the individual infection a diagnostically characteristic appearance.

CHAPTER V

TOXIN AND ANTITOXIN

THE REACTION BETWEEN TOXIN AND ANTITOXIN (EHRlich'S ANALYSIS)

THE TOXIN-ANTITOXIN REACTION

WHEN Behring and his collaborators, Kitasato and Wernicke, had definitely shown that the cell-free blood serum of animals immunized with tetanus and diphtheria toxins respectively possessed the power to protect other animals of the same and different species against the poisons, it became of the utmost importance to determine, if possible, the mechanism by which the "antitoxic" effect was attained. The earlier opinion, expressed by Behring himself, held that in all probability the toxin was directly injured or destroyed by the action of the antitoxic serum. That this assumption was incorrect was soon demonstrated by the experiments of Roux and Vaillard¹ and by those of Buchner.² The work of the former investigators showed that the mixtures of tetanus toxin and antitoxin, measured in such proportions that they were harmless for normal guinea pigs, could still be found toxic for animals weakened by preliminary inoculation with other bacteria. Buchner claimed in analogous experiments that similar mixtures, harmless for mice, could still show toxicity for guinea pigs. He inferred from this that the nature of the cell reactions of different animal species influenced the antitoxic effect. Both investigations led the workers to conclude that the protective action of antitoxin was not due to a direct effect upon the poison but was potent by acting upon the tissue cells of the animal by protecting these from subsequent harm by the toxin. Their conception implied an indirect protective function on the part of the antitoxin, not due to any direct reaction between it and the poison.

That this explanation, too, was faulty was made evident by a number of investigations which took advantage of the peculiar differences in resistance to temperature between certain toxins and their specific antitoxins.

¹ Roux and Vaillard. *Ann. de l'Inst. Past.*, 1894.

² Buchner. *Münch. med. Woch.*, p. 427, 1893.

In 1894 Calmette³ and Physalix and Bertrand⁴ had independently succeeded in obtaining an antitoxin against snake poison. In the course of further study of these bodies Calmette⁵ determined that the venoms of certain varieties of snakes, the naja and cobra, would remain potent even when subjected to 100° C. for a very short time. In contrast to this the antitoxins to these poisons were destroyed at much lower temperatures. Now when mixtures of the two substances, so proportioned that their injection into animals was innocuous, were heated to 68° C. for considerable periods, toxic properties again became evident, a demonstration that the toxin had not been destroyed, but had remained neutral only in the presence of the intact antitoxins. These experiments were confirmed by Wassermann,⁶ who found that similar conditions prevailed in the combination between pyocyaneus toxin and antitoxin.

The filtration experiments of Martin and Cherry⁷ are not convincing since they may be taken as indicating either neutralization or toxin destruction. These workers subjected mixtures of snake poison and its specific antitoxin to filtration through gelatin filters, under pressure. Under the experimental conditions thus established the presumably smaller toxin molecule was allowed to pass through the filter while the larger antitoxin molecule was held back. They showed that if filtered soon after the ingredients have been put together most of the toxin still passes through, but that, as this interval is prolonged, less and less comes through, presumably because of the union of the smaller toxin to the larger antitoxin molecule. The chief value of these experiments lies in their proof of the element of time as an important factor in the toxin-antitoxin union.

In his experiments on snake venom just recorded, Calmette interpreted the restitution of toxicity after the heating of neutral mixtures of cobra neurotoxin and its antitoxin as evidence "qu'il ne s'était pas formé aucune combinaison de ces deux substances ou que la combinaison réalisée était, au moins, très instable." Later experiments of Martin and Cherry seemed for a time to contradict this conclusion. Observations by them, analogous to those of Calmette, but carried out with the poison of an Australian snake, seemed to indicate that when the toxin and antitoxin were allowed to remain together for a sufficiently long time no restitution of toxicity could be obtained by heating. Apparently the application of heat to such mixtures merely prevented the further union of antitoxin with any toxin that was not yet bound at the time that the heat was applied. Accord-

³ Calmette. *Compt. rend. de la soc. de biol.*, 1894.

⁴ Physalix and Bertrand. *Compt. rend. de la soc. de biol.*, 1894.

⁵ Calmette. *Ann. Past.*, 1895.

⁶ Wassermann. *Zeitschr. f. Hyg.*, 22, 1896.

⁷ Martin and Cherry. *Proc. of the Royal Soc.*, Vol. 63, 1898.

ingly Morgenroth⁸ again examined these relations and found that the addition of a small amount of hydrochloric acid to mixtures of snake poison and the antitoxin resulted in the dissociation of their union. To mixtures of the venom lysin and its antitoxin, neutralized and even overneutralized so that they were perfectly innocuous to susceptible animals he added hydrochloric acid until the total concentration amounted to N/18. By this method a toxin-HCl modification was produced which was dissociated from its union with the antitoxin and was extremely resistant to heat. In such a mixture of toxin and antitoxin to which hydrochloric acid had been added, heating at 100° C. in a water bath for 30 minutes destroyed the thermolabile antitoxin and, after neutralization, undiminished toxic properties could again be demonstrated by animal inoculation.

These researches and other similar ones of Morgenroth, then, form a satisfactory confirmation of the original experiments of Calmette and seem to show, beyond possibility of contradiction, that the inhibition of harmful properties of any true toxin, after mixture with its antitoxin, does not depend upon toxin destruction. But while Calmette interpreted the facts as pointing toward a failure of union of the two substances, Morgenroth's work is not incompatible with the conception of a neutralization of one by the other in the chemical sense. These experiments of Morgenroth are of great theoretical importance moreover in that they have shown that dissociation of a toxin-antitoxin complex can occur.

The nature of such neutralizations in regard to quantitative relations, speed of action, and relative concentrations, becomes apparent partly from experiments like those mentioned above, but more especially from those carried out by Ehrlich with ricin and antiricin, experiments which were primarily planned to demonstrate that the reaction between a toxin and its antibody is a direct one, not dependent upon intervention of the body cells, as at first supposed.

It had been shown by Kobert and Stillmarck that ricin, the powerfully poisonous principle of *Ricinus communis* (castor oil bean) would agglutinate the red blood cells of a number of animals. Ehrlich recognized from the beginning how closely analogous the neutralization of ricin by antiricin was to that of diphtheria toxin by its antitoxin. The former reaction furnished him with a simple method of test tube experimentation in that the agglutinating effects of ricin upon rabbits' corpuscles could be directly inhibited by the preliminary addition of antiricin. A visible reaction was thus available, which, of course, excluded absolutely the participation of the tissue cells in the antigen-antibody neutralization, and in which careful quantitative measurements were possible.

Ehrlich⁹ determined by means of this method that the neutral-

⁸ Morgenroth. *Berl. kl. Woch.*, No. 50, 1905, p. 1550.

⁹ Ehrlich. *Fortschr. d. Med.*, Vol. 15, p. 41, 1897.

ization was accelerated by moderate heat and by concentration of the reagents and, most important of all, that the reaction followed roughly the law of multiple proportions, characteristics, all of them, which were entirely analogous to chemical reactions in general. When he added 0.3, 0.5, 0.75, 0.1, etc., cubic centimeters of serum from a ricin-immune goat to constant quantities of ricin, and then added rabbit cells, the hemagglutinating properties of the ricin were inhibited in direct proportion to the amount of antiricin mixed with it. And his test tube experiments were further found to represent with much accuracy the occurrences which took place within the animal body. For, similar mixtures injected into mice were toxic in direct proportion to the balance of ricin and antiricin established in the injected material.

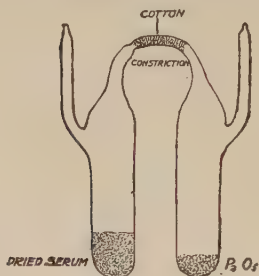
Although the views of Ehrlich and his followers have great importance in connection with the union of antigens and their antibodies in general, these ideas were worked out by him most elaborately in connection with his efforts to arrive at a practicable and accurate method of establishing a standard of strength for diphtheria antitoxin, and it is essential that we consider this work in detail.

The earlier attempts to standardize diphtheria antitoxin by the use of living cultures (Roux and Behring) were soon abandoned, since it was found that the accurate establishment of fixed lethal doses of the culture was not possible. When the facts, just recorded, concerning the interaction and quantitative relations of the soluble toxins and their respective antitoxins came to light, Behring introduced the standardization of the curative sera by the use of toxins, both in the case of tetanus and in that of diphtheria. In order to do this consistently he established for diphtheria poison an arbitrary toxin unit which he defined as the amount of any given diphtheria filtrate sufficient to cause death in a guinea pig of 250 grams, and, borrowing the terms from chemical nomenclature, he designated as a "normal" diphtheria poison one which contained 100 such units in one cubic centimeter. (D T N, M250 = diphtheria toxin normal, Meerschweinchen 250 grams.)

Together with Ehrlich, Behring then established an antitoxin unit (I-E, Immunitäts Einheit). They designated as a "normal" antitoxic serum one "which contained in one cubic centimeter one antitoxic unit" (I-E), and state further, "of this serum 0.1 c. c. neutralizes 1 c. c. of the Behring normal toxin." (Conf. Madsen in "Kraus u. Levaditi Handbuch," II, p. 94.) Alterations were subsequently made in this scale of standards and Ehrlich later designated as an antitoxin unit a quantity of an antitoxin which completely neutralized 100 lethal doses (for guinea pigs of 250 grams) of a poison at that time in his possession. The unit of diphtheria antitoxin at present in use therefore may be defined as a quantity of serum sufficient to protect a guinea pig of 250 grams

against 100 times the fatal dose (M L D, minima dosis lethalis) of toxin. Since the methods of antitoxin standardization employed at present in the United States were worked out by Rosenau¹⁰ along the lines of Ehrlich's method, and the standard is based on the one introduced by Ehrlich, the antitoxin unit as employed in this country is identical with the one spoken of in the following paragraphs.

In measuring the neutralizing value of antitoxin for toxin, then, since both substances are chemically unknown and no purely chemical indicator of neutralization is available, it was necessary to select a susceptible animal by means of which excess of toxin, in mixtures of the two, could be detected. As the standard test animal guinea pigs of 250 grams were chosen, and improvements in the methods of measurement were introduced, in that the toxin and antitoxin, instead of being separately injected as heretofore, were mixed, allowed to stand for 15 to 30 minutes, and then injected together subcutaneously.



TUBE FOR THE PRESERVATION OF THE STANDARD ANTITOXIN.

Taken from Rosenau, U. S. Hygienic Laboratory Bulletin, No. 21, 1905, p. 53.

By means of this technique Ehrlich set out to examine a large number of toxins and their antibodies and obtained results which, aside from their practical value, have had an important influence upon the development of the knowledge of antigen-antibody reactions.

These investigations were considerably complicated by the fact that neither the diphtheria toxin nor the antitoxin is very stable and deterioration occurs unless special methods of preservation are employed. Since the antitoxin, however, is much less unstable than the toxin, the former is employed in order to preserve the standard, and is preserved in sealed U tubes (see Figure) with anhydrous phosphoric acid. Kept in this way, in black, light-proof boxes, and at low temperature, it may be preserved for months without appreciable loss of value and may be renewed by accurate comparative measurements from time to time. This is carried out for the United States, at the present time, by the Government Hygienic Laboratories at Washington.

Preservation of the toxin is much more difficult, and it is in connection with the investigation of the instability of the toxin that Ehrlich gained his first insight into the nature of the reaction. He measured, in a number of toxic filtrates, the minimal lethal dose for guinea pigs of 250 grams, establishing a time limit for death in order to obtain more accurate comparisons. He designated as the

¹⁰ Rosenau. U. S. P. H. & Mar. Hosp. Service, *Hygienic Laboratory Bull.*, 21, April, 1905.

new M L D or "T" (that is: toxic unit) the quantity of toxin which will kill a guinea pig of the designated weight in from 4 to 5 days. He then determined for a number of poisons the exact quantity just neutralized by one antitoxin unit, calling this quantity L_0 . (L meaning Limes or threshold.)

It is clear that in judging of complete neutralization of a quantity of toxin by antitoxin, there may be a strong subjective element, since any very slight excess of toxin may cause unimportant local reactions such as edema or small hemorrhages, which could escape the attention of one observer while being noticed and recorded by another. In order therefore to exclude definitely all subjective features from such experimentation, Ehrlich now established another toxin value— L_+ dose ("Limes death"—now, for convenience, written L_+)—which eliminated all possible variations of personal perception. He designated by this symbol that quantity of toxin which not only neutralized one antitoxin unit but included enough toxin, in excess of this, to give the result of one free toxin unit, that is, to cause death in 4 to 5 days in a guinea pig of 250 grams. Since the three values just defined form the basis of Ehrlich's experiments as well as that of all practical diphtheria serum standardizations we will briefly restate them for the sake of clearness.

Thus:

M L D or "T" = the amount of toxin which, subcutaneously injected, causes death in a 250-gram guinea pig in from 4 to 5 days.

L_0 = the amount of toxin which is just neutralized by one antitoxin unit so that no trace of reaction, local or otherwise, ensues

and

L_+ = that amount of toxin which will cause death in 4 to 5 days in a guinea pig of 250 grams if injected together with one antitoxin unit.

It will further clarify the meaning of these terms to examine experimental protocols which show how these values are determined.

Thus in the following:

I. Injections of toxin

- (1) .005 c. c.—guinea pig lives.
- (2) .009 c. c.—guinea pig dies in 6 days.
- (3) .01 c. c.—guinea pig dies in 4 days.
- (4) .02 c. c.—guinea pig dies in 2 days.
- .01 = M L D or T.

II. 1 Antitoxin unit + .19 toxin = late paralysis.

1 Antitoxin unit + .20 toxin = sometimes late paralysis and sometimes acute death.

1 Antitoxin unit + .21 toxin = death fourth day.

1 Antitoxin unit + .22 toxin = death in 2 to 3 days.

.21 = L_+ dose.

- III. 1 Antitoxin unit + .14 c. c. toxin = no reaction.
 1 Antitoxin unit + .15 c. c. toxin = no reaction.
 1 Antitoxin unit + .16 c. c. toxin = slight congestion about point of injection, scarcely visible.
 1 Antitoxin unit + .17 c. c. toxin = apparent reaction at site.
 1 Antitoxin unit + .18 c. c. toxin = edema at site.
 $L_0 = .16$.¹¹

In determining these values with a large number of toxins Ehrlich discovered the curious fact that, although there was a rapid and extensive diminution of toxicity in every toxic filtrate in the course of time, there was no corresponding alteration in the L_0 amount. In other words, although more and more of the toxic broth was necessary to kill a guinea pig of standard weight in the required time, the amount of the same broth which neutralized one antitoxin unit remained approximately the same.¹²

In seeking an explanation for this apparent paradox, Ehrlich concluded that we must assume that the toxin is complexly constructed, consisting of a toxophore and a haptophore group. Assuming that chemical union between the toxin and the antitoxin (or, in disease, the body cell) takes place, it is by means of the haptophore group that such union is brought about. The toxophore group, however, is the element by which toxic action is exerted after union by the haptophore group has been accomplished. It would be conceivable, therefore, that in deteriorating in toxicity the toxin might undergo alterations in the toxophore group only, its haptophore group, and, therefore, its antitoxin neutralizing properties remaining intact. Such modified toxins, modified only as to the toxophore groups, Ehrlich now refers to as "toxoids."

In the production of diphtheria toxins for practical purposes it has been found advisable to allow them to "season," that is to stand for prolonged periods until they have reached a state of "equili-

¹¹ II and III are taken from the article by Rosenau, *P. H. & M. H. S., Hyg. Lab. Bulletin*, 21, 1905.

¹² This statement plainly contradicts the definition of a toxin unit; i. e., the amount which neutralizes 100 toxin units and often leads to confusion among students or others who are unfamiliar with this subject. It should be borne in mind that, while the definition of an antitoxin unit is the one accepted when Ehrlich first arbitrarily established it, the antitoxin unit, as at present in use, is really an amount of antitoxin standardized against L_0 quantities of toxin, this last value again obtained by measurement against the original unit. It represents a neutralization value equal to the original one, but may protect the guinea pig against 85, 110, 130, etc. (variable) toxin units, according to the constitution of the particular toxic filtrate employed in the experiment. Indeed, if, in the following pages, the reasoning of Ehrlich is consistently adhered to, our definition of an antitoxin unit should be: The amount of antitoxin which contains 200 binding affinities for toxin. This will become clearer as the following paragraphs are read.

brium" at which the conversion of toxin to toxoids has been reduced to a minimum and the change of relationship between L_0 and "T" or M L D has practically ceased to go on. From the very beginning of the growth of the culture in the incubator the process of toxoid formation has probably occurred, and even freshly prepared toxic filtrates therefore are not pure "toxins," especially since the conversion of toxin to toxoid seems to diminish in velocity as time goes on.

Now in spite of the presence of such alteration products, in comparing the values L_0 and L_+ of any given toxin preparation, one would naturally suppose that L_+ minus L_0 should be equal to one M L D, or the quantity just sufficient to kill a guinea pig of 250 grams in 4 to 5 days. For we have seen that L_0 just neutralizes one antitoxin unit while L_+ is the quantity which, in addition to such neutralizing power, has an excess of toxin equal in action to one minimal lethal dose. This, however, is not the case. Let us illustrate this by a concrete case. One of Ehrlich's toxins on measurement showed a minimal lethal dose or M L D of 0.0025 c. c.

The L_+ dose of this was .25
while The L_0 dose of this was .125

The difference was .125 or 50 M L D instead of 1 M L D as we would suppose.

Stated in words, this measurement means that after neutralizing completely one antitoxin unit with the toxic filtrates, in order to obtain death in a guinea pig in 4 days with such a mixture, it was necessary to add, beyond the neutralizing quantity, 50 M L D, or again as much as was necessary for neutralization.

This last relation is merely coincidence, since it might have been 30 or 40 or 60 M L D just as well. The important point is the fact that more than 1 M L D was necessary, and by this fact Ehrlich was led to resort to an assumption which forms one of the basic principles of many of his explanations for serum phenomena, namely, the assumption of differences in combining avidity or affinity.

As applied to the present problem he reasoned as follows:

It is conceivable that the toxoids resulting from deterioration of toxin might possess three different degrees of affinity for the antitoxin. They might have a stronger, an equal, or a lesser affinity than the toxin itself. If their affinity for antitoxin were equal to that of toxin they would, of course, not influence the L_+ dose itself; if stronger than toxin their influence would be so exerted that toxin would be forced out of combination with antitoxin, giving place to the toxoid, and the effect would be the opposite from that experimentally observed. If, however, their affinity for antitoxin were weaker than that of toxin each additional toxin unit added to the L_0 dose would unite with antitoxin, replacing a corresponding quan-

tity of the toxoid of weaker affinity. In consequence, as far as the poisonous properties of the mixture are concerned, the addition of toxin would not render the neutral mixture poisonous for guinea pigs until the toxoids had been completely displaced from union with antitoxin. Finally, after all the antitoxin had been bound to unchanged toxin, further addition would then result in the presence of free toxin and poisonous properties would again appear in the mixture. Ehrlich at first spoke of the toxoids possessing weaker affinity for antitoxin than the toxin itself as "epitoxoids." This conception can be rendered clear by the following example:

In the case cited above we had

$$T \text{ or } M L D = 0.0025 \text{ c. c.}$$

$$L_+ = 0.25 \text{ c. c.}$$

$$L_0 = 0.125 \text{ c. c.}$$

$$\text{The difference} = 0.125 = 50 M L D.$$

Supposing that the toxoids (epitoxoids) present in the mixture possessed an affinity for antitoxin less than that of toxin, the following conditions might obtain:

$$151 \text{ toxin-antitoxin} + 49 \text{ epitoxoid-antitoxin} = L_0.$$

Add 1 M L D or T and we have:

$$152 \text{ toxin-antitoxin} + 48 \text{ epitoxoid-antitoxin} + 1 \text{ epitoxoid free.}$$

Add 2 M L D or T and we have:

$$153 \text{ toxin-antitoxin} + 47 \text{ epitoxoid-antitoxin} + 2 \text{ epitoxoid free until, finally, adding } 50 T, \text{ we get:}$$

$$200 \text{ toxin-antitoxin} + 49 \text{ epitoxoid free} + 1 \text{ toxin free} = L_+.^{13}$$

Later experience led Ehrlich to abandon the opinion that the epitoxoids were deterioration products of the toxin. He found that the relation between L_0 and L_+ which we have just outlined, was demonstrable in the same way, in freshly prepared toxin filtrates, in which there had been little time for toxoid formation. He further

¹³ An example identical in significance with the one just given, but somewhat simpler in its arithmetical conditions, is here added for the sake of permitting no possibility of unclarity. This example is taken from Ehrlich's own work.

$$T = .01 \text{ c. c. of the toxin bouillon.}$$

$$L_+ \text{ (neutral. of antitoxin unit yet killing 1 pig)} = 2.01 \text{ c. c. or } 201 T.$$

$$L_0 \text{ (complete neutral. of 1 antitoxin unit)} = 1 \text{ c. c. or } 100 T.$$

$$\text{Difference} = 1.01 \text{ c. c. or } 101 T.$$

$$100 \text{ toxin-antitoxin} + 100 \text{ epitoxoid antitoxin} = L_0;$$

Add 1 T, and we have:

$$101 \text{ toxin-antitoxin} + 99 \text{ epitoxoid-antitoxin} + 1 \text{ epitoxoid free;}$$

Add 101 T, and we have:

$$200 \text{ toxin-antitoxin} + 100 \text{ epitoxoid free} + 1 T \text{ free} = L_+.$$

noticed that, even after deterioration had occurred to a considerable extent, and the amount necessary to kill a guinea pig had been much increased (the number of fatal doses in L_0 constantly decreasing as toxoids replaced toxin), the L_+ nevertheless remained unchanged. This, he held, could mean one thing only. The elements present in toxic broth which possessed a weaker affinity for antitoxin than the toxin itself, namely, the epitoxoids, were present from the very beginning and were probably separate and primary secretion products of the diphtheria bacilli, remaining relatively stable and constant as the broth was preserved. In order to avoid confusion, therefore, he now referred to the "epitoxoids" as "toxons"—to preclude their confusion with the other toxoids or true toxin deterioration products. These toxons possess, according to Ehrlich, a "haptophore" group identical with that of the toxin, but have a different toxophore group. For there is reason to believe that the toxon, lacking the power of causing acute death, gives rise to slow emaciation and paralysis, finally killing after a subacute or chronic course. Thus, in the tabulation just preceding, we have seen that the toxic broth added to neutral mixtures of toxin and antitoxin (containing the L_0 dose), does not give rise to the acutely toxic effect of one M L D or T until an amount has been added which considerably exceeds one toxin unit. This, we explained, by Ehrlich's reasoning, on the supposition that "epitoxoids" or "toxons" are displaced from their union with antitoxin, giving place to toxin and becoming free. Such toxin-antitoxin mixtures—in which the amount of toxin broth ranges between the L_0 and the L_+ doses—therefore, contain no free toxin units but contain varying amounts of free toxon. An injection into guinea pigs is not followed by acute death in these cases, but leads with considerable regularity to emaciation, paralysis, and death after a long incubation period.

It has been objected to this, as we shall see, that the slow poisoning produced by such mixtures is due, not to a qualitatively different poison but to the presence of minute quantities of free toxin too small to produce acute death, yet sufficient to produce this gradual injury. This Dreyer and Madsen¹⁴ tried to disprove by experiments in which they prepared antitoxin-toxin mixtures so balanced that they possessed the toxon effect, and of these mixtures injected increasing multiples. In no case did they succeed in producing acute death even when the amount injected had been multiplied to such an extent that free toxin, if present, must have asserted itself. The same workers were able to show that the injection of these mixtures, in which free toxons were assumed to be present, produced immunity against toxin, thus indicating the similarity of the haptophore group of toxin and toxon—a conception which will

¹⁴ Dreyer and Madsen. *Zeitschr. f. Hyg.*, Vol. 37, 1901.

become more and more clear as we consider the "Side-Chain Theory" which Ehrlich evolved as a result of his toxin analysis.

Ehrlich had thus elicited facts which seemed to him to indicate the presence of three qualitatively different substances in toxic filtrates of diphtheria cultures. Two of these, the toxin and the toxon, were present, he assumed, in freshly prepared filtrates, as independent primary secretion products of the bacilli, the toxin an acute poison, the toxon a substance with slower and qualitatively different poisonous effects. Both of them, toxin and toxon, possessing similar haptophore groups, could unite with antitoxin and neutralize it, but the affinity of toxon for antitoxin was weaker than that of toxin. For this reason toxin could displace toxon from its union with antitoxin, this accounting for the discrepancy between the L_+ and the L_0 doses. The third class of substances, the toxoids, were deterioration products of toxin, the deterioration implying an alteration in the toxophore group only, the haptophore group remaining the same.

It is plain from this reasoning that Ehrlich's conception implies complete analogy between chemical reactions in general and the neutralization of toxin by antitoxin. Accordingly it is but another step in the same direction to speculate concerning the actual relations of valency existing between the two substances. It seemed to Ehrlich that there were many reasons for assuming that the union between toxin and antitoxin occurred in proportions of 200 to 1; that is, just as the formula for water is H_2O , that of toxin-antitoxin combinations would be "Toxin₂₀₀Antitoxin."

The considerations on which this opinion was based were as follows: In examining a large series of toxic filtrates, Ehrlich,¹⁵ as well as Madsen, had found that the number of toxin units ("T" or M L D) necessary to neutralize one antitoxin unit (that is, the number of toxin units contained in the L_0 dose) corresponded, with great regularity, to multiples of 100. Values of 25, 33, 50, 100, etc., recurred again and again. This indicated that the deterioration of the toxin into toxoids followed a certain regularity of progression and seemed to justify the assumption that the absolute binding power possessed by antitoxin was represented by a valency corresponding to a multiple of 100. Since the number of toxin units contained in an L_0 dose rarely fell below, and usually above 100, the valency could not be less than 100. On the other hand, repeated measurements of L_0 and L_+ doses never showed as many as 200 toxin units. Ehrlich's own highest value was 133, and the highest ever obtained by any one was a measurement by Madsen of 160. Now considering the fact that no toxin is "pure" but that, in every case, it contains admixtures of toxoid and toxon, the values 133 or 160 cannot represent all the valencies of an antitoxin unit. They represent only that part of the antitoxin unit which is neutralized

¹⁵ Ehrlich. *Deutsche med. Woch.*, No. 38, 1898, Vol. 24.

by the "toxin," as measurable upon guinea pigs, a certain fraction of antitoxin being united to toxoid or toxon. It is likely, therefore, as Ehrlich reasoned, that, being higher than 100, and in an obviously impure condition approaching but never reaching 200, the valency of antitoxin for toxin was just 200. The correctness of this surmise seemed rendered more probable by Ehrlich's further studies, since analysis of the ingredients of various toxic filtrates, that is, the determination of the relative contents of toxin, toxoids, and toxon, appeared to show constantly definite relations to the valency 200.

The method by which Ehrlich carried out these subsequent studies is spoken of as the method of "Partial Absorption." In principle it represents a reversal of his earlier methods of measurement. In these he had titrated various amounts of toxin broth against a constant quantity (one unit) of antitoxin, establishing the L_+ and L_0 values. In the method of Partial Absorption, on the other hand, he measured varying fractions of an antitoxin unit against a constant amount of toxin, employing for this a previously determined L_+ and L_0 dose. A measurement carried out in this way is shown in the following tabulation in which, at the same time, there is indicated how many valencies each antitoxin fraction represents, on the basis of an assumed total of 200 for each unit.¹⁶

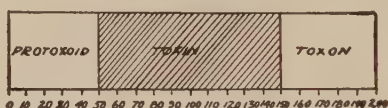
| | | | |
|------------------------------|---------------|---------------|--------------|
| 0 antix. unit representing | 0 valency | + L_+ = 85 | free T units |
| .1 antix. unit representing | 20 valencies | + L_+ = 85 | free T units |
| .25 antix. unit representing | 50 valencies | + L_+ = 60 | free T units |
| .8 antix. unit representing | 160 valencies | + L_+ = 10 | free T units |
| .9 antix. unit representing | 180 valencies | + L_+ = 3.5 | free T units |
| 1.0 antix. unit representing | 200 valencies | + L_+ = 1 | free T unit |

It is immediately evident in this table, as it would be evident in any other citation of similar measurements, that there is no regularity in the progress of neutralization; or, in other words, that addition of a definite fraction of antitoxin does not remove the arithmetically corresponding amount of toxic properties from the L_+ dose. The first 0.1 unit of antitoxin in this table has removed no free toxin whatever. The addition of the next 0.15 of an antitoxin unit, representing 30 valencies, has removed $\frac{2}{3}$ or $\frac{5}{7}$ of the total toxicity. Throughout the scale there is not the regular progression of neutralization, multiple by multiple, which would be expected if antitoxin could be titrated against a pure toxin. This, according to Ehrlich, is due to the presence of various toxoids which possess varying affinities for the antitoxin molecule. The first 0.1 of a unit added, in this case, does not diminish the toxicity of the mixture because it is bound by "protoxoids" which possess a higher affinity for antitoxin than the

¹⁶ This measurement is taken from one cited by Ehrlich in *Deutsche med. Woch.*, No. 38, 1898, Vol. 24, and is taken literally except for the first value of 1/10 antitoxin unit, which is inserted to illustrate the formation of protoxoids.

toxin itself. Next are bound the toxins themselves together with varying amounts of "syntoxoids" which possess the same affinity as toxin. Finally there are left the toxons which possess a lesser affinity than toxins or toxoids, and therefore again have the discrepancy between the L_0 and L_+ dose. Ehrlich utilizes this method in order to determine the composition of the constituents of any given toxic filtrate and expresses the results in the so-called "toxin spectra."

The construction of these spectra and the principles underlying the measurements on which they are based are very clearly illustrated by Madsen,¹⁷ from whose article the following type spectra are taken:¹⁸

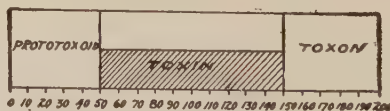


TOXIN SPECTRUM AFTER MADSEN, *Ann. de l'Inst. Past.*, Vol. 13, 1899, p. 57.

This figure represents the diphtheria filtrate composed of 50 valencies of protoxoid, 100 toxin and 50 toxon equivalents. The measurements in this case may be represented by the following tabulation:

| | | |
|-------------|---------------------------------|------------------|
| $L_0 + 1$ | antitox. unit = 200 valencies = | 0 lethal dose |
| $L_0 + .75$ | antitox. unit = 150 valencies = | 0 lethal dose |
| $L_0 + .25$ | antitox. unit = 50 valencies = | 100 lethal doses |
| $L_0 + 0$ | antitox. unit = 0 valency = | 100 lethal doses |

The following diagram, also from Madsen, represents the same poison after it had deteriorated to $\frac{1}{2}$ its toxic power. L_0 , therefore, would contain only 50 toxic doses.



AFTER MADSEN, *Ibid.*, p. 578.

The measurements corresponding to this table are as follows:

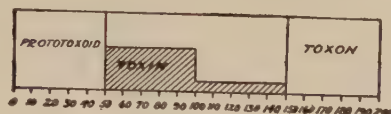
| | | |
|--------------|---------------------------------|---------------|
| $L_0 + 1.$ | antitox. unit = 200 valencies = | 0 lethal dose |
| $L_0 + .75$ | antitox. unit = 150 valencies = | 0 lethal dose |
| $L_0 + .745$ | antitox. unit = 149 valencies = | 0 lethal dose |
| $L_0 + .74$ | antitox. unit = 148 valencies = | 1 lethal dose |
| | etc. until | |

¹⁷ Madsen. *Ann Past.*, Vol. 13, 1899, p. 576.

¹⁸ We have chosen to illustrate the principles of the toxin spectrum from the article of Madsen rather than from Ehrlich's original work, because the former presents this difficult phase of the subject in a hypothetical toxin of extremely simple structure. Some of Ehrlich's spectra constructed from actual measurements may be found in *Deutsche med. Woch.*, No. 38, 1898.

$L_0 + .25$ antitox. unit = 50 valencies = 50 lethal doses
 $L_0 + 0$ antitox. unit = 0 valency = 50 lethal doses

The following spectrum, the third in Madsen's article, represents the same toxin described in the preceding spectrum but at a later period, at which considerable further deterioration had taken place. The L_0 dose now contained but 30 M L D or, in other words, the amount of toxin contained in the L_0 dose was sufficient to kill 30 guinea pigs only.



AFTER MADSEN, *Ibid.*, p. 579.

Madsen's description of the method in which this spectrum is constructed is the following:

| | |
|--|----------------|
| $L_0 + \frac{200}{200}$ of an antitoxin unit kills | 0 guinea pig |
| $L_0 + \frac{190}{200}$ of an antitoxin unit kills | 0 guinea pig |
| $L_0 + \frac{180}{200}$ of an antitoxin unit kills | 0 guinea pig |
| $L_0 + \frac{170}{200}$ of an antitoxin unit kills | 1 guinea pig |
| $L_0 + \frac{160}{200}$ of an antitoxin unit kills | 2 guinea pigs |
| $L_0 + \frac{150}{200}$ of an antitoxin unit kills | 3 guinea pigs |
| $L_0 + \frac{140}{200}$ of an antitoxin unit kills | 5 guinea pigs |
| $L_0 + \frac{130}{200}$ of an antitoxin unit kills | 5 guinea pigs |
| $L_0 + \frac{120}{200}$ of an antitoxin unit kills | 6 guinea pigs |
| $L_0 + \frac{110}{200}$ of an antitoxin unit kills | 6 guinea pigs |
| $L_0 + \frac{100}{200}$ of an antitoxin unit kills | 7 guinea pigs |
| $L_0 + \frac{90}{200}$ of an antitoxin unit kills | 10 guinea pigs |
| $L_0 + \frac{80}{200}$ of an antitoxin unit kills | 30 guinea pigs |
| $L_0 + \frac{70}{200}$ of an antitoxin unit kills | 30 guinea pigs |

The amount of toxon has remained the same in spite of deterioration. As less and less antitoxin is added, between the values of $\frac{150}{200}$ and $\frac{100}{200}$ of an antitoxin unit, there are now liberated only 5 fatal doses of the toxin. It is in this zone that deterioration has taken place, since, in the preceding spectrum, the difference between the addition of $\frac{150}{200}$ and $\frac{100}{200}$ of an antitoxin unit represented 25 fatal doses for guinea pigs. When in this last spectrum the amount of antitoxin is gradually reduced from 100 valencies to 50 valencies 25 fatal doses are liberated, a quantity corresponding to the similar zone in the preceding spectrum. Thus in this particular zone of the spectrum no change has taken place. The same is true of the protoxoid zone.

It is unnecessary to cite a larger number of such measurements in this place, since the ones given sufficiently illustrate the methods

and the conclusions drawn from them. As a result of such experiments Ehrlich concludes:

I. That the diphtheria bacillus produces primarily two kinds of substances (a) toxin, (b) toxon, both of which bind the antibody.

II. The toxins (and perhaps also the toxons) may deteriorate and be modified into secondary substances (toxoids) which may be distinguished by their different degrees of affinity for antitoxin.

III. This classification does not exhaust all possible complications, since each subdivision of toxin consists apparently of equal parts of two different modifications which are similar to each other in their relation to antitoxin but differ in varying resistance to influences of deterioration. A more complete analysis of these conditions may be found, together with a series of illustrative spectra, in Ehrlich's article in the *Deutsche med. Wochenschr.*, Sept., 1898, which has been quoted above.

The complex deductions arrived at by Ehrlich are largely dependent, as we have seen, upon strict adherence to the analogy between the toxin-antitoxin reactions and those occurring between strong acids and strong bases. In such cases there is a complete reaction, in which chemical change ceases only when there has been a complete neutralization of one by the other. If, for instance, we mix molecular equivalent amounts of H_2SO_4 and NaOH , an apparently complete change into Na_2SO_4 and H_2O occurs:



The reverse process does not seem to take place, and if traces of uncombined H_2SO_4 and NaOH are present, as may be theoretically assumed, they are so slight in amount that they are not demonstrable. There are, however, many chemical reactions in which the process is not a complete one, in that the chemical change does not proceed until the reagents are completely used up. Reaction in these cases ceases when an equilibrium is reached at which there are present definite amounts of the reaction products and of the original substances at the same time.¹⁹

This occurs when a weak acid is added to a weak base. In such cases the reaction is incomplete and reversible and, together with the neutralization products, both free acid and free base may be present. The conditions are best explained by citing an example of a reversible reaction which is commonly given in text-books of physical chemistry, namely, the reaction between ethyl-alcohol and acetic acid. (Our citation is taken from Philip's "Physical Chemistry," London, Arnold, 1910): "When one gram mol. of ethyl alcohol is added to one gram mol. of acetic acid, a reaction takes place which results in

¹⁹ See Cohn. "Vorträge f. Ärzte über Physik. Chem.," Engelmann, Leipzig, 1901.

the formation of ethyl acetate and water; the reaction, however, is incomplete and stops at an equilibrium point at which the reaction mixture contains $\frac{1}{3}$ gram mol. alcohol, $\frac{1}{3}$ gram mol. acid, $\frac{2}{3}$ gram mol. ethyl acetate, and $\frac{2}{3}$ gram mol. water. If, on the other hand, 1 gram mol. of ethyl acetate is mixed with 1 gram mol. of water, a reaction sets in which results in the formation of ethyl alcohol and acetic acid. This change likewise stops in equilibrium at a point at which the composition of the reaction mixture is the same as that already stated." The reaction is thus reversible and may be written:



Another example somewhat simpler and more easily brought into analogy with the toxin-antitoxin reaction is that of the dissociation of phosphorus pentachlorid into phosphorus bichlorid and chlorin (see Alexander Smith, "General Chemistry," Century Company, N. Y., 1911, p. 181).

Here the reaction takes place:



Chemical equilibrium is reached when the reaction speed is the same in both directions, and there will be present, at equilibrium, PCl_3 , Cl_2 , and PCl_5 . Now the "Law of Mass Action" (Guldberg & Waage) states that reaction goes on at a velocity proportionate to the concentration of the reacting molecules. It is plain, therefore, that at the point at which the reaction takes place with equal velocities in both directions, that is, at the equilibrium point, a very definite relation of molecular concentrations must obtain, and this relation can be expressed as a formula. For the example given above this may be written as follows:

$$\frac{\text{Conc. PCl}_3 \times \text{Conc. Cl}_2}{\text{Conc. PCl}_5} = K \text{ (constant)}$$

This formula is expressed in words by Alexander Smith as follows: "If we change the amount of the pentachlorid placed in the vessel, or if we use amounts of chlorin and trichlorid which are not equivalent, the numerical value at equilibrium of each concentration will, of course, be different, but the product of the concentrations of trichlorid and chlorin, divided by the concentration of the pentachlorid, will always give the same numerical value for (K), the constant, at the same temperature."

Now to return to the application of these facts to the neutralization of toxin by antitoxin, if the reaction is one analogous to that of a strong acid and alkali, as cited above in the case of H_2SO_4 and

NaOH, we would expect a complete neutralization of one by the other, multiple for multiple, and the explanation of Ehrlich based on the assumption of different toxin constituents, of varying affinities, and different pharmacological effects, is the only one which will account for the experimental results. Assuming, however, that the reaction is one analogous to that taking place between a weak acid and a weak base—such as boric acid and ammonia—we have an entirely different state of affairs. For here the reaction goes on to a point of equilibrium, and in mixtures containing equivalent amounts of the weak acid and the base there will be present the reaction products and also small amounts of unbound free acid and free base. And according to the law of "Mass Action," the quantities of free acid and base present will depend entirely on the masses of the reagents put together. Thus, for each particular mixture of the two, different quantities of the original substances will be found uncombined, and, furthermore, the gradual addition of one to the other will not have a neutralizing value in exact proportion to the amount added. Were the toxin-antitoxin reaction analogous to such chemical systems, then we could assume that every mixture of the two substances, whatever the relative amounts, would contain not only the united toxin-antitoxin molecule, but also varying amounts of dissociated free toxin and free antitoxin, the quantities of each depending, according to the law of mass action, upon the molecular concentrations obtaining in the individual mixture. This, indeed, is the conception of toxin-antitoxin union formulated by Arrhenius and Madsen.

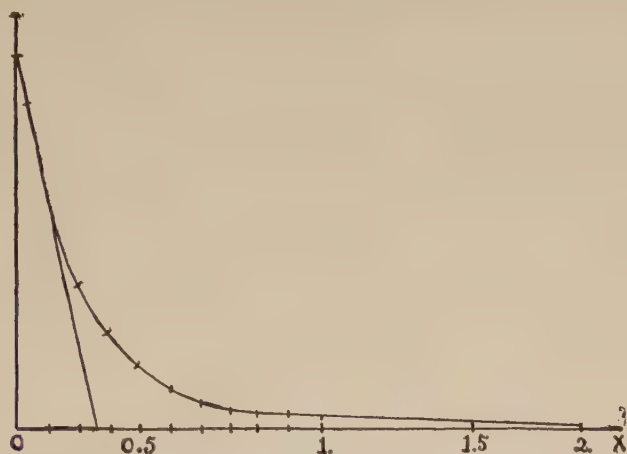
Arrhenius and Madsen,^{20 21} bearing in mind these conditions, made comparative studies of the neutralization of tetanolyisin by its antilysin on the one hand, and that of ammonia by boric acid on the other. Ammonia, like most bases, is a hemolytic agent, while boric acid, unlike stronger acids, has no hemolyzing properties. For this reason, in mixtures of the two, the toxicity is proportional to the concentration of free ammonia (though, as Arrhenius states, "a correction must be made for the lowering action of the ammonium salt, as indicated by experiments on this action"). Because the reaction between boric acid and ammonia is reversible, that is, the salt formed is dissociated by the hydrolytic effect of the water, there is always present a slight amount of free ammonia even if the largest possible quantities (to saturation) of boric acid are added. (See Arrhenius, "Immunochem.," p. 174.) The curve of toxicity indeed descends as more boric acid is added, but never reaches 0.

By a modification of the formula expressing the law of Mass Action, Arrhenius and Madsen could calculate the amount of free

²⁰ Arrhenius and Madsen. *Zeitschr. f. physik. Chem.*, 44, 1903, and *Festschrift Kopenhagen Serum Instit.*, 1902.

²¹ Arrhenius. "Immunochemistry," Macmillan, N. Y., 1907.

ammonia present in a series of mixtures in which increasing quantities of boric acid were added to a constant quantity of ammonia, and



CURVE REPRESENTING THE NEUTRALIZATION OF TETANOLYSIN BY DIFFERENT QUANTITIES OF ANTITOXIN.

Taken from Arrhenius, "Immunochemistry," Macmillan, 1907, p. 175.

the values so obtained corresponded with much accuracy to those resulting from measurements of toxicity upon red blood cells. The following table taken from Arrhenius and Madsen illustrates this:

TOXICITY (q) OF 0.1 N. NH_3 (1 EQUIVALENT) WITH n EQUIVALENTS OF BORIC ACID. (Taken from Arrhenius, *loc. cit.* p. 176.)

| n = Equivalents of boric acid added | Quantity of free ammonia—i. e., toxicity—observed | q = Ammonia toxicity calculated from formula | Δq obs. |
|---|---|--|-----------------|
| 0 | 100 | (100) | |
| 0.17 | 85 | 79 | 15 |
| 0.33 | 69 | 64 | 16 |
| 0.67 | 43 | 42 | 26:2 = 13 |
| 1 | 25 | 27 | 18:2 = 9 |
| 1.33 | 20 | 18 | 5:2 |
| 1.67 | 13 | 13 | 7:2 |
| 2 | 10 | 10 | 3:2 |

Here, in the last column, there is indicated the proportion of toxicity which is neutralized by the successive addition of $\frac{1}{6}$ of an equivalent of boric acid. The first additions lower it to a degree proportionate to the amount of acid added; the next additions neutralize it to a much slighter degree, and, as further additions are

made, each successive one possesses progressively less relative neutralizing power than the preceding.

This, it is plain, is closely analogous to the phenomena observed by Ehrlich in his "Partial Absorption" method, and Arrhenius concludes that the two phenomena, toxin-antitoxin and boric acid-ammonia neutralization, are closely analogous. His point of view is further strengthened by his experiments with tetanolysin and its specific antibody, in which he constructed a curve similar to that given for boric acid, derived a formula and found that the observed and the calculated values closely coincided for various mixtures of the two. He claims, in consequence, that the phenomena observed by Ehrlich should not be interpreted as due to "partial toxins"—toxoids or toxons, but dependent rather upon the presence of varying quantities of free toxin dissociated from union with antitoxin because of the reversibility of the union.

The opinions of Arrhenius and Madsen are not generally accepted. It is in the first place doubtful whether substances like toxin and antitoxin, which, as far as we know their chemical nature at all, belong to the class of substances spoken of as colloids, can be regarded as subject to the laws of Mass Action in their reactions.

Nernst²² has criticized Arrhenius' deductions chiefly on the basis of their assumption of the reversibility of the union of toxin and antitoxin, since reversible reactions between colloids, though not at all inconceivable, have so far not been definitely shown. Furthermore, as Nernst states, if complete reversibility of such reactions were possible it would be hard to understand how antitoxin can protect the animal against the actions of toxin.

Another point of view concerning the toxin-antitoxin union which has been gaining ground especially through the work of Landsteiner and his pupils, is that of Bordet.²³ Bordet expresses his views in the following way:

I. When one mixes with a certain quantity of toxin an amount of antitoxin which is insufficient to produce a complete neutralization, the molecules of antitoxin are not taken up by a definite fraction of the toxin molecules, satisfying the affinities of these entirely while other units remain intact; on the contrary, the antitoxin molecules distribute themselves equally upon all the toxin molecules present, and these are therefore, all of them, partially saturated, and lose proportionately a part of their initial toxicity. One could say that there is an attenuation of the toxin since there is a formation of a less poisonous complex.

II. The symptoms of poisoning produced by such a complex injected into animals or placed in contact with sensitive cells cannot be

²² Cited from Landsteiner in "Kolle u. Wassermann Handbuch," 2d Ed., Vol. 5.

²³ Bordet. *Ann. de l'Inst. Past.*, Vol. 17, 1903.

identical with those which would be produced by a fully saturated mixture of toxin and antitoxin, or by intact toxin.

III. Between these two extremes, free toxin and entirely neutralized toxin, one can imagine many transitions, progressive stages of attenuation. Every time that one mixes toxin and antitoxin in the same way one attains the same degree of attenuation.

Briefly put, this means that Bordet estimates toxin-antitoxin combinations of different degrees of toxicity as representing different stages in the completeness of the saturation of the individual toxin units. When 10 parts of toxin are added to 1 part of antitoxin, the result, according to him, would not be such that 1 part is neutralized by 1 part of antitoxin, leaving 9 parts of toxin free. He assumes rather that each unit of toxin is attenuated by the absorption of $\frac{1}{10}$ th of a part of antitoxin. He compares this process to the action of iodine upon starch. Starch can absorb variable quantities of iodine and, according to the amount taken up, is colored slightly or deeply blue. This mode of action is common to most staining processes. The substance that is stained fixes varying quantities of coloring matter and the coloring matter does not limit itself to a definite fraction of the substance stained but distributes itself equally to the material, coloring it slightly or deeply, in its entirety, according to the relative amount of color added. We will see later that there are many reasons for regarding other antigen-antibody combinations as following similar laws of proportion.

Bordet and others speak of this point of view as the "Absorption Theory," and Biltz, in studying this point of view by physical methods, comes to the conclusion that the observed figures of the quantitative relations between toxin and antitoxin in the process of neutralization are fairly consistent with the values to be expected if the process were actually an absorption phenomenon.

A curious occurrence which seems to bring the toxin-antitoxin reactions close to colloidal reactions in general is that which is known as the "Danysz²⁴ Effect" or as the "Bordet²⁵-Danysz Phenomenon." Danysz discovered that when ricin or diphtheria toxin were brought into contact with their homologous antibodies the degree of neutralization depended upon the manner in which the two were put together. When the toxin was added to the antitoxin in two fractions, a considerable time being allowed to elapse between the additions, the final mixture was much more toxic than when the total amount was added at once. In other words, although both mixtures contained exactly the same quantities of the two reacting substances, nevertheless the amount of toxin left free varied in the two cases, according to the speed with which they had been put to-

²⁴ Danysz. *Ann. de l'Inst. Past.*, Vol. 16, 1902.

²⁵ Bordet. *Ann. de l'Inst. Past.*, Vol. 17, 1903.

gether. This was confirmed in 1904 by von Dungern²⁶ for diphtheria toxin, and Craw²⁷ was able to observe it in the case of megatheriolysin and its antilysins.

Von Dungern interpreted this in the sense of Ehrlich, by assuming it to be due to what he calls "epitoxonoids." This epitoxonoid he assumes to be a constituent of toxic broth, which has still less affinity for antitoxin than the toxon. It can combine with diphtheria antitoxin, but not until all the true toxin is bound. However, when it is once united with diphtheria antitoxin it is not very easily displaced from the union, especially when a considerable time has elapsed since the union. Therefore, he thinks, when the toxin is added to the antitoxin in two fractions, this epitoxonoid is bound and keeps the toxin, which is added later, out of combination. Whereas if the toxic broth is added as a whole, it is the epitoxonoid which is left unbound. This explanation of von Dungern's may be looked upon as an ingenious refinement of the reasoning introduced by Ehrlich into this field. As a matter of fact reactions similar to the Danyz phenomena have been very commonly observed in the reactions between various colloids.

THE SIDE-CHAIN THEORY

Mechanism of Antibody Formation

The discovery of antitoxins in the blood serum of toxin-immune animals by Behring and his collaborators furnished a point of new departure for the investigation of the phenomena of immunity, and Ehrlich's work upon the nature of the reaction between toxin and antitoxin, both in vitro and in the animal body, firmly established that the protective effect of the latter was one of direct neutralization, and not, as at first supposed, one of toxin destruction or of indirect influence through the mediation of the body cell. As we have seen, moreover, it was quickly noted that these reactions were strictly specific in that an antitoxin produced with any one of the known toxins reacted solely with this one to the exclusion of all others. All these facts were of the utmost practical importance and gave hope of ultimate extensive therapeutic application, a hope which has, in part, been realized.

The physiological mechanism by which these phenomena were brought about, however, was, and is, to a great extent still, a mystery, and a most extensive and painstaking series of researches has occupied itself with its elucidation.

²⁶ Von Dungern. *Deutsche med. Woch.*, 30, 1904.

²⁷ Craw. *Jour. Hyg.*, Vol. 7, 1907.

When we consider the invariable production of a specific antitoxin in response to the treatment of an animal with a toxin it is but natural that Buchner and others should have at first assumed that the antitoxin is, in each case, a product obtained by the action of the body tissues from the toxin itself. While difficult to refute at a time when little was known of the laws of antitoxin production and of quantitative relationships, such an assumption is entirely untenable in the light of more recent knowledge. We now know that such a simple conversion of toxin into antitoxin cannot explain the phenomenon because the amount of antitoxin incited in the immunized animal is out of all proportion great in comparison with the amount of toxin injected. Thus Knorr²⁸ has found that 100,000 units of antitoxin may be produced by the injection of the toxin equivalent of one unit. Moreover the discovery by Salomonsen and Madsen²⁹ that pilocarpin injections will increase the amount of antitoxin produced by an animal distinctly pointed to the likelihood of the participation of the general physiological activities of an immunized subject in the production of antibodies. Unquestionable proof of this was also brought by the experiments of Roux and Vaillard,³⁰ in which antitoxin production in immunized animals continued even after the entire volume of blood had been removed by repeated bleeding. This observation points distinctly to the direct secretion of antibodies by the tissue cell, in the nature of what has been termed by Roux³¹ an "internal secretion." And it is this activity of the body cell in the production of antibodies which forms the fundamental premise from which the now classical "Side-Chain Theory" of Ehrlich takes its departure.

In order to approach this theory logically it will be of advantage to consider briefly the general subject of the assimilation of food-stuffs and other substances distributed by the circulation to the cells of the animal body. For, as Ehrlich has expressed it, "The Reactions of Immunity, after all, represent only a repetition of the processes of normal metabolism, and their apparently wonderful adjustment to new conditions is only another phase of 'Uralter Protoplasma Weisheit.'" ³² It is impossible to conceive the nutrition of body cells without assuming that the assimilable nutritive substances come into physical and, eventually, chemical relationship with the protoplasm of the nourished cell. Considering the large variety of substances which may thus be brought into contact with cells in the course of normal and abnormal metabolism, the body cell,

²⁸ Knorr. *Münch. med. Woch.*, 1898, pp. 321, 362.

²⁹ Salomonsen and Madsen. *Ann. de l'Inst. Past.*, Vol. 12, 1898.

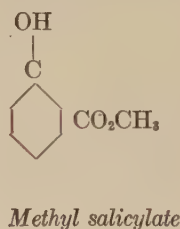
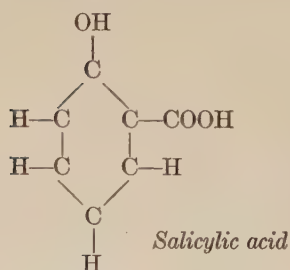
³⁰ Roux and Vaillard. *Ann. de l'Inst. Past.*, Vol. 7, 1893.

³¹ Roux. Ref. in *Semaine Medicale*, 1899.

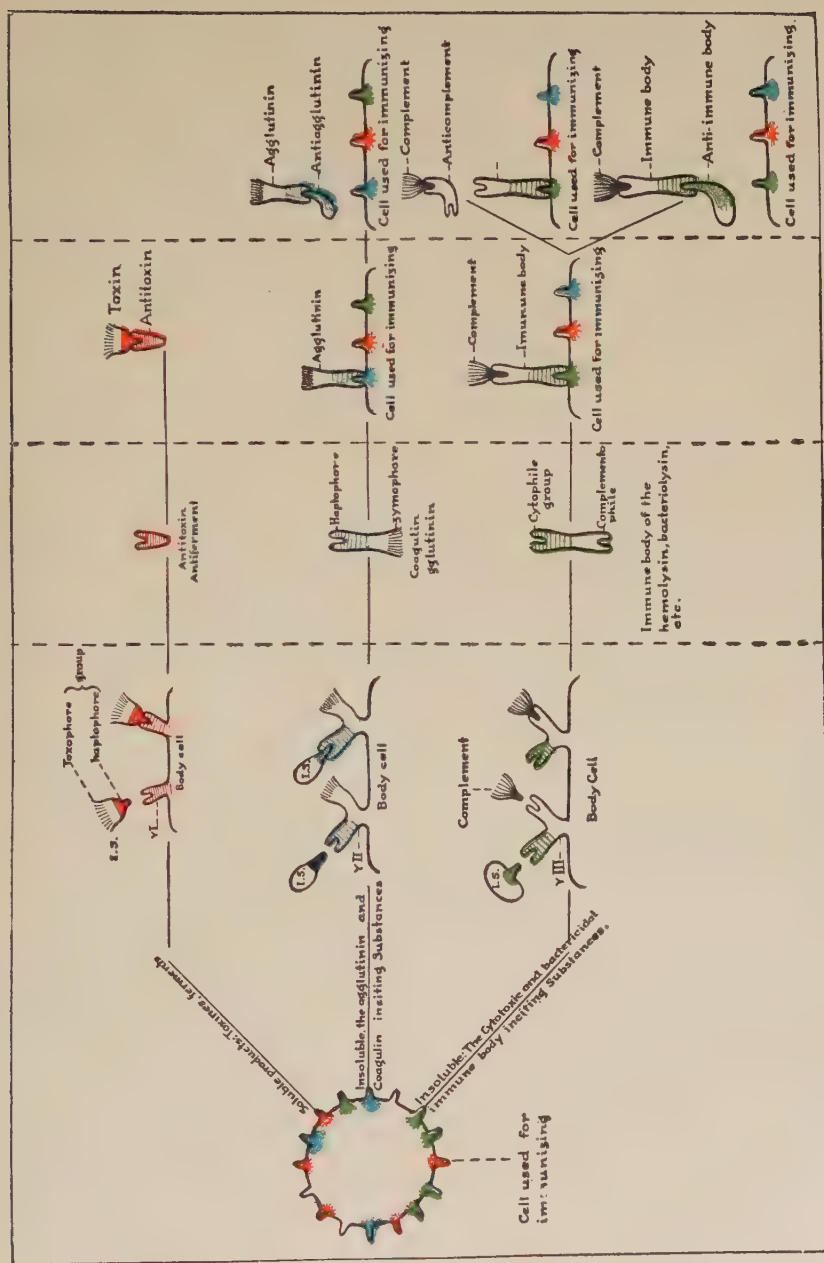
³² Ehrlich. Introduction to "Gesammelten Arbeiten," Berlin, Hirschwald, 1904.

chemically considered as a complex of enormous molecules, must possess a correspondingly great variety of atom groups, by means of which it can unite with these substances to assimilate them and make them a part of its own protoplasm. In order to enter into similar relationship with toxins and other antigens, then, it is only logical to suppose that the cell, in the same way, unites chemically with the antigenic substance, and either assimilates it without sustaining harm, as in the case of non-poisonous complexes, or is injured in the process, as in the case of the poisons.

The living cell, from this point of view, is conceived as consisting of a central chemical nucleus, the "Leistungskern," more or less stable, in that the specialized tissue function is dependent upon it, and a manifold variety of "side chains," or atom groups by means of which it can enter into relationship with the nutritive and other materials carried to it by the body fluids. The latter term, "side-chains," is taken from the nomenclature of chemistry, and, although the analogy is a loose one, it serves satisfactorily to elucidate Ehrlich's meaning. Thus we may conceive the "Leistungskern" as the central carbon ring of any compound of the Benzol series, as, for instance, in salicylic acid in which the hydrogen atoms, the hydroxyl,



and the acid radicles represent "side chains." By means of the latter the compound can enter into relation with other substances, as, for instance, with CH_3 in the formation of methyl salicylate. Graphically, though this analogy formulates Ehrlich's fundamental conception, it must not be taken as too literally representing the existing conditions, since, in actual metabolic interchange, there is an infinite variety of possible "side-chain" groups; for we are dealing with an enormous number of assimilable substances, most of them of chemically unknown constitution. The cell, therefore, is looked upon as an active chemical complex, retaining its own peculiar functional characteristics by reason of the "Leistungskern," but constantly getting rid of waste products and entering into new union with extraneous materials by virtue of its "side chains." These side chains, because of their "receiving" function, are spoken of by Ehrlich as cell "receptors."



THE STRUCTURE OF CELL-RECEPTORS AND IMMUNE BODIES, ACCORDING TO EHRLICH'S CONCEPTION.
(After Aschoff.)

That the chemical structure of certain bodies determines their ability to enter into relation with cell derivatives such as enzymes is, of course, a fact well established by experiment and explains the specific action of bacterial and other ferments upon certain substances to the exclusion of others. Thus Pasteur noted the fact that bacterial ferments could decompose dextrorotatory tartaric acid while they did not affect the levorotatory variety, and Emil Fischer³³ showed that only those carbohydrates possessing 6 and 9 carbon atoms were subject to fermentation by yeasts, and of these only the ones belonging to the "d" series, observations which, by demonstrating the relationship between these active agents of extracellular digestion, and the stereochemical configuration of the molecules acted upon, lend much support to the logic of Ehrlich's contentions.

Moreover, the recent experiments upon the growth of tissues in plasma outside of the animal body in which cartilage cells produce cartilage, kidney cells, etc., have shown that, given the same nutritive materials, the cells themselves must command a certain selective power in the choice of these materials, which can only depend upon a specific element in the structure of the cell receptors. As Fischer has expressed it for fermentation, the ferment must possess an atom group which fits into some group of the fermentable substance as a "key does into a lock," an analogy which is equally applicable to Ehrlich's conception of the relation of the "side chain" to a nutritive molecule.

Now the toxins and other antigens are, all of them, so far as we know, complex chemical substances, derivatives of animal and vegetable cells, and, for this reason, should have much in common with the materials available for nutrition. It is not strange, therefore, that, coming into contact with the cells of the body during the accidents of disease or other abnormal conditions, they should find receptors by means of which they can combine with the cell. Under the ordinary conditions of nutrition a suitable particle taken up by the cell in this way would be assimilated and the receptor either freed for further use or regenerated for the further absorption of similar substances, by virtue of a mechanism delicately coördinated to the needs of cell-nutrition. In the case of the absorption of substances belonging to the class of antigens, however, foreign proteins difficult of assimilation, or of toxins even directly harmful, the receptors occupied by these substances are rendered useless to the cell, and, if the cell continues to live, must be regenerated. If the degree of poisoning or the amount of other antigen introduced has been extremely slight, this regeneration may possibly take place, as in the course of nutritive processes, without further disturbance. If, however, the amounts of antigen are greater than this, or are repeat-

³³ See Oppenheimer, "Die Fermente," Vol. 1.

edly thrust upon the cell, the process of regeneration may be not only sufficient to compensate for the loss of the eliminated receptors, but may follow the general law of overcompensation, formulated by Weigert, and receptors of the variety occupied by the antigen are produced in excessive number.

Here again Ehrlich has called analogy to his aid, and has taken his conception of "overcompensation" from the well-known phenomena of pathological anatomy where, for instance, in the restoration of cellular elements after injury, there is often an overproduction of granulation tissue, far beyond the needs of simple healing.

Thus the restitution of cell receptors, if sufficiently stimulated by large quantities or repeated administration of the antigen, far exceeds the quantity normal to the cell, and may proceed to such a degree that the cell, becoming as it were "top-heavy" with these elements, sloughs them off into the surrounding lymph and blood, where they circulate as free receptors. These free receptors then, having specific affinity and combining power for the antigen which incited their production, unite with subsequently introduced antigen in the blood stream, diverting it from the cells themselves, and, in the case of the variety of antigens spoken of as toxins, this union with the free receptors in the blood stream would serve to protect the cells from harm, exerting thereby an antitoxic action.

The antibodies appearing in the blood of immunized animals, therefore, represent atom complexes, normally parts of the body cells and concerned in the metabolic processes, but now produced in excess and extruded into the body fluids under the influence of the stimulation of immunization. The very substances, as Behring has put it, which make possible the poisoning of the cell by the toxins become protective when, detached from the cell, they circulate in the blood. Thus the theory, beside explaining the causes leading to antibody formation, offers a plausible reason for the relatively strict specificity observed in antibody-antigen reactions.

Formulated in direct connection with the investigations upon toxins and antitoxins, the side-chain theory has been extended by Ehrlich and his associates to all known phases of antibody-antigen reactions. The differences in the nature and complexity of various antigens would naturally necessitate variation in the receptors capable of assimilating them, and these receptors, appearing subsequently in the blood as antibodies, must, of necessity, differ from each other. On this basis Ehrlich has conceived of three main varieties or "orders" of receptors or "haptines," as he calls them. Of these the simplest are those of the first order which attach to the *toxins*, and by over-regeneration appear in the blood stream as *antitoxins*. Those of the second order, adapted to the assimilation of more formidable protein molecules, are, of necessity, of greater structural complexity, appearing in immunized animals as the *agglutinins* and *precip-*

itins, while those of the third order, dependent upon the coöperation of alexin or complement, for proper functioning, appear as the *cytotoxins* or *lysins*. The detailed structure of these various haptines will be discussed in connection with other considerations dealing with their special reactions.

Limiting ourselves, for the present, to a broad consideration of the theory as a whole, it may be briefly recapitulated as follows: Toxins or other antigens, in order to exert any influence upon the animal body, must enter into chemical relationship with the cells. This they do by virtue of union with chemical units or atom groups of the cells, spoken of as "side chains." These side chains or receptors, thrown out of function by this union, and necessary for the metabolic processes of the cell, are regenerated, and under the influence of repetition of this process are produced in excess, to such a degree that they are eventually thrown off by the cells and enter the circulation as antibodies. Thus far the theory, comparing the union of antigen with cells to the processes of nutrition, is eminently logical and likely, necessitating the assumption of over-regeneration as the only criterion not directly amenable to experimental proof.

That the antigen can be bound by the body cells has been variously shown in a large number of investigations, some of which have been reviewed in our section on the action of bacterial poisons. We have there seen that Dönitz demonstrated the rapid disappearance of tetanus and diphtheria toxins from the circulation of susceptible animals, and that conversely Metchnikoff showed that the poison may persist unabsorbed and unchanged for weeks and months in the blood of such insusceptible animals as the turtle and the lizard, facts which furnish indirect evidence of the absorption of the toxins by the body cells. More direct evidence has, of course, been possible in the test tube experiments with hemolytic and other cell poisons where a directly specific combination between antigen and antibody has been easily demonstrable. Thus, in his earlier experiments with spider poison, Sachs was able to show that rabbit erythrocytes, which are sensitive to the poison, could absorb it out of solution, while dog and other corpuscles, which were insusceptible to the poison, did not bind or absorb it. This can be easily demonstrated for many antigens and antibodies and may be accepted as a fact.

This point established, and repeatedly confirmed, and the origin of antitoxins from the cells of the body having been rendered likely by the experiments of Salomonsen and Madsen, and by those of Roux and Vaillard just cited, it would follow, by the theory of Ehrlich, that we should find the site of antibody production in the very cells which possessed specific affinity (receptors) for the antigen. This question has been variously investigated, chiefly in the case of the toxins and antitoxins, since this phase of the subject is most easily amenable to experiment. It will be remembered also that Wasser-

mann and Takaki discovered that emulsions of the tissue of the central nervous system of rabbits and guinea pigs, shown by Meyer and Ransom and others to be the special points of attack for tetanus toxin, possessed the power of neutralizing this poison in vitro, while emulsions of spleen kidney and other organs had no such effect. They assumed from this that the poison was fixed by cell receptors, antecedents of antitoxin in the sense of Ehrlich. Kempner^{34 35} made similar observations with botulinus toxin and further confirmation has been derived from experiments like those of Blumenthal,³⁶ who found that the toxin was neutralized by the brain tissue of susceptible animals but showed conversely that the brain substance of the chicken, an animal but slightly susceptible to tetanus, possessed little or no neutralizing power. Similar results were obtained by Metchnikoff in the cases just cited.

The great importance of these experiments lies not only in showing that body cells may absorb the toxins, but that there is direct relationship between the susceptibility of tissue and the toxin-binding properties. Furthermore the facts demonstrated by Metchnikoff that no antitoxin was produced by those animals (turtle, lizard) in which the tissues had no power of fixing poison and which are consequently insusceptible, furnish powerful evidence in favor of Ehrlich's view.

It becomes of great importance, therefore, to determine whether in the case of the fixation of tetanus toxin by the brain cells the union between cell and toxin is a specific and chemical one comparable in every way to the union of toxin with antitoxin.

Metchnikoff, in spite of his results in the experiments just cited, objected to this interpretation on the ground that although the brain emulsion of a guinea pig neutralized tetanus toxin *in vitro*, the injection of the toxin into such an animal, subdurally, produced the disease. This can hardly be regarded as a valid argument against Wassermann's interpretation, since the very premises of the Ehrlich theory require that these neutralizing elements, when still attached to the living cell, as "sessile" receptors, are the cause of the poisoning, since they serve to "unlock" the cell to the entrance of the toxin. Similar objections on the part of Metchnikoff³⁷ were based on some of his own experiments, as well as on those of Courmont^{38 39} and Doyen, in which it was found that the poison disappears but slowly (in 2 to 3 months) from the circulation of frogs, and the brain cells show hardly any toxin neutralization in vitro, whereas these animals

³⁴ Kempner and Pollak. *Deutsche med. Woch.*, 1897, No. 23, p. 505.

³⁵ Kempner and Shepilewsky. *Zeitschr. f. Hyg.*, Vol. 36, 1901, p. 1.

³⁶ Blumenthal. *Deutsche med. Woch.*, 1898, No. 12, p. 185.

³⁷ Metchnikoff. *Ann. de l'Inst. Past.*, Vol. 12, 1898.

³⁸ Courmont and Doyen. *Arch. de Physiol.*, 1893.

³⁹ Courmont and Doyen. *Compt. rend. de la soc. de biol.*, 1893.

can be rendered tetanic if they are warmed to 25° to 30° C. Further work, however, by these authors as well as by Morgenroth⁴⁰ has satisfactorily cleared up this difficulty. As a matter of fact, tetanus poison disappears more rapidly (that is, is bound by the cells more rapidly) from the circulation of frogs, if the frogs are warmed to 30° C. or more. Furthermore, if the toxin is injected into these animals, and they are kept at low temperatures, no disease results, but if they are then warmed up to the temperature stated, they gradually succumb to the disease. Morgenroth has shown that the apparently anomalous behavior of frogs in this respect is actually a question of temperature. At low temperatures the poison is bound, though with extreme slowness, but the toxophore group of the toxin does not functionate. When the animals are warmed, not only does the binding proceed more rapidly, but the toxophore group becomes active. He thus not only has answered Metchnikoff's objections to Ehrlich's theory on this ground, but has furnished an additional indirect confirmation of the dual constitution of toxin, that is, its constitution of a haptophore and a toxophore atom group, suggested by Ehrlich in his diphtheria-toxin analysis.

There is apparently, then, a strong absorption of tetanus toxin by the brain and nervous tissue of all animals which are susceptible to the poison, an absorption which amounts, as we have seen, to neutralization, the brain emulsion acting like antitoxin when mixed with the toxin before injection, as in Wassermann's and Takaki's experiments.

A serious objection has been brought, however, to the assumption that this binding can be identified in its nature with the similar binding of toxin by antitoxin, and a number of authors have claimed that the binding by the brain is not a binding by specific receptors, but an accidental property due to the presence of some fortuitous fixing substance in the central nervous system. Besredka⁴¹ showed, for instance, that the brain of susceptible animals could bind much more toxin than it could actually neutralize, and that, if antitoxin was added to a brain emulsion previously saturated with the toxin, the toxin is removed from its combination with the brain cells and these again regain their original absorbing property. These experiments would seem to point to a difference, especially in regard to affinity and firmness of union between the nature of the combination between toxin and brain emulsion on the one hand, and toxin and antitoxin on the other. This, of course, would prove a serious obstacle to the interpretation of the binding of toxin by susceptible cells in the sense of Ehrlich, as depending as it were upon union with specific receptors, or, as they might be termed, "sessile" antitoxin. Moreover, to strengthen such objections to this point of view, the work of Land-

⁴⁰ Morgenroth. *Arch. internat. de Pharm.*, Vol. 7, 1900, pp. 265-272.

⁴¹ Besredka. *Ann. de l'Inst. Past.*, Vol. 17, 1903, p. 138.

steiner and v. Eisler⁴² has brought out the fact that extraction of brain tissue with ether materially reduces its toxin-binding powers by removing fatty or lipoidal substances, such as cholesterin and lecithin. And it has indeed been confirmed that lipoids can possess, in many instances, binding properties not only for toxins but for other forms of antibodies. On the basis that at least a part of the toxin absorption by brain emulsions depends upon such lipoidal fixation, the results of Besredka are readily explained, but were this the sole cause of toxin fixation by these tissues it would indeed be difficult to interpret the phenomenon, with Wassermann and Takaki, in support of Ehrlich's theory. For, without going into further refinements, the fact of the probable proteid, certainly not lipoidal, nature of the antitoxins, discussed in a previous section, would alone serve to distinguish the two modes of toxin fixation.

However, a number of facts have been ascertained which show that, although the lipoids play some part in the antitoxic action of brain cells, they do not by any means account for the entire process. In the first place it is found that the heating of brain emulsions almost completely removes their power to bind the toxin, while no such reduction of the fixative property follows the heating of lipoids like cholesterin or lecithin. The experiments of Marie and Tiffeneau⁴³ have done much to clear up the confusion regarding this point. They determined that the "lipoidal binding" constituted only about one-tenth of the total binding power of the brain emulsions, by showing in the first place that only one-tenth of the total was left after heating, and that all but one-tenth could be destroyed by subjecting the tissue to the action of proteolytic enzymes. It appears from this that a large part, at any rate, of the toxin fixation of the brain tissues is dependent upon substances of an albuminous nature, a smaller but definite part being dependent upon fixation by lipoids, a phenomenon entirely apart from the former in underlying principles. This would, it seems, both justify the original interpretation of Wassermann and still explain the apparently contradictory results of Besredka and others.

⁴² Landsteiner and v. Eisler. *Centralbl. f. Bakt.*, Vol. 39, 1905.

⁴³ Marie and Tiffeneau. *Ann. de l'Inst. Past.*, Vol. 22, pp. 289 and 644, 1908.

CHAPTER VI

THE BACTERICIDAL PROPERTIES OF BLOOD SERUM, CYTOLYSIS, AND SENSITIZATION

IN spite of the profound physiological alteration of the animal body which is implied by the acquisition of immunity against any particular infection, we have seen that no anatomical or histological changes in the organs and tissues accompany such alteration. The same is true of the difference between animals of different species, in which the most marked variation in resistance against any given infection is inexplicable on the basis of structural or microscopic characteristics in the organs. We have mentioned briefly the attempts that have been made to discover chemical and physical changes or differences to account for such conditions and have seen that the attention of investigators was soon attracted to the blood.

A possible relationship between the blood and the defence of the body against infection had been foreshadowed by observations made long before the days of bacteriological knowledge. As early as 1792, John Hunter, in his "Treatise on the Blood, Inflammation and Gun-shot Wounds," had noted that the blood did not decompose as readily as other putrescible material, and a century later, during the period of great interest in the living nature of fermentation and putrefaction, Traube (1874) expressed the opinion that blood could destroy bacteria. Similar observations were made by Lister and by Grohman¹ but no experimental work aimed at this point was carried on until 1886, when the subject was taken up by Nuttall,² von Fodor,³ and Flügge, and a little later by Buchner.⁴ These authors, working with defibrinated blood, peptone blood, and blood serum, showed that such substances all exerted a definitely measurable destructive influence upon bacteria, and Nuttall, later confirmed by Buchner, further found that this bactericidal power was weakened on standing, and could be rapidly destroyed by heating to 60° C.

Their method of procedure consisted in the planting of controlled amounts of various bacteria in measured quantities of blood and,

¹ Grohman. Quoted from Adami, "Principles of Pathology," Vol. 1, p. 497.

² Nuttall. *Zeitschr. f. Hyg.*, 4, 1888.

³ Von Fodor. *Deutsche med. Woch.*, 1887.

⁴ Buchner. *Centralbl. f. Bakt.*, Vol. 5, 1889.

after several hours at 37° C., pouring plates and thus determining the numbers of surviving organisms. The fact of bactericidal power established, there was, of course, much early difference of opinion as to the mechanism responsible for the destruction of the bacteria, and a number of simple explanations were suggested which, though entirely refuted at the present time, still possess considerable interest in showing the stages of development through which the conceptions of the mechanism of immunity have progressed.

These early theories were formulated chiefly upon the underlying thought that the animal body was primarily passive in its relation to the invading micro-organisms, and that the disappearance of bacteria in the body fluids was due to the existence of a chemically or physically unfavorable environment which prevented their multiplication and therefore induced gradual mortality among them. Thus Billroth⁵ believed that bacteria could thrive in the body only after a preceding putrefactive change had prepared a favorable pabulum. Others attempted to discover a relation between the degree of alkalinity of the blood serum and the destruction of bacteria. This argument was soon refuted by the experiments of Buchner, who showed conclusively that the bactericidal power of serum was not reduced by the neutralization of its natural alkalinity with weak acetic acid.

Another theory which has been kept alive until the present day by Baumgarten,⁶ and in favor of which much has been written by Fischer, is the so-called "Osmotic" explanation. The basis of this conception is the observation that vegetable and other cells, which are in themselves delicate osmotic systems, undergo changes when they are placed into fluids of different osmotic tension.⁷ Thus, of course, cells of all kinds may be destroyed by being placed in distilled water on the one hand, or in hypertonic salt solution on the other. The point of view of Baumgarten, as explained in a recent edition of his "Text-book of Bacteriology," is the following: The bacterial (or blood) cell, like all cells, is surrounded by a semi-permeable membrane. Under ordinary conditions, this membrane permits the passage of certain substances which must enter and leave the cell in the course of normal metabolism. When the bacteria are placed in a specific bacteriolytic serum there is a chemical union between the antibody and the cell membrane, and the latter is, in consequence, injured. The result of the injury is that now the cell becomes permeable for salts and other substances to which it was impermeable before, and there are consequent swelling and in-

⁵ Billroth. Quoted from Sauerbeck, "Die Krise in der Immunitätsforschung," Klinkhardt, Leipzig, 1909.

⁶ Baumgarten. "Lehrbuch der pathogenen Mikroorg.," Hirzel, Leipzig, 1911.

⁷ See also Pfeiffer's "Pflanzen Physiologie."

creased intracellular pressure. This, in turn, brings about the extrusion from the cell of proteins and other ordinarily non-diffusible substances, and destruction of the cell results. This explanation is practically an adaptation of the earlier more primitive osmotic theories to the facts subsequently discovered. It stands in direct contradiction to the prevailing opinion that the process of bacteriolysis and cytolysis in general is an enzymotic process, brought about by the injury of the cell by specific substances comparable to digestive ferments. Interesting though the suggestion of Baumgarten is, it can hardly receive more than casual attention given it for the sake of completeness, since careful experimental work by von Lingelsheim⁸ has shown definitely that altered salt contents of serum do not exercise the effect upon bacteriolysis which we would be entitled to expect from Baumgarten's reasoning.

In explanation of the natural immunity possessed by many animals against various infections, Baumgarten has offered another explanation which, like the preceding, we may classify, in agreement with Sauerbeck,⁹ with the "passive" theories. This theory, which he calls his "Assimilation Theory," assumes that the bacteria do not find suitable food material in the tissues and fluids of certain animals, and, since bacteria do not have to be killed to be eliminated, but may be checked merely by their inability to grow and multiply, they must soon succumb in surroundings in which they find no suitable foodstuffs. This point of view approaches somewhat the earlier exhaustion theory of Pasteur, which has been mentioned in another place.¹⁰

In contrast to these "Passive" theories of immunity are the now prevailing and well-founded opinions that the resistance of the animal body against bacterial invasion is not a mere fortuitous result of chemical and physical conditions encountered by the infectious agents, but is rather the result of the struggle against the invasion by active forces of the body cells and fluids. The part played by the cells had already been emphasized by Metchnikoff and his school when the discovery of the bactericidal power of the normal blood was made. The study of the antibacterial powers of the blood now introduced a new element which became the basis of the so-called "humoral" theories. In the prolonged controversies waged, with great astuteness and experimental skill, between the adherents of these two schools, most of the facts which we possess regarding immunity were discovered, and it is only within recent years that we

⁸ Von Lingelsheim. *Zeitschr. f. Hyg.*, Vol. 37, 1901.

⁹ Sauerbeck. "Die Krise in der Immunitätsforschung," Klinckhardt, Leipzig, 1909.

¹⁰ The influence of foodstuffs, temperature, and other environmental conditions upon natural immunity has been discussed in an earlier section.

have obtained information which has made possible a correlation between these two main paths of thought.

The humoral theory was conceived by Buchner, as the first important theoretical result of Nuttall's discovery. Buchner, as we have seen, confirmed the observations of Nuttall both as to the primary fact of the bactericidal power of the fresh normal blood and as to the unstable nature of this bactericidal property. He looked upon the antibacterial power as depending upon a constituent of the fresh blood plasma, which he named "*alexin*" (protective substance), and which he believed to be comparable to a proteolytic enzyme. The action of this alexin was conceived as potent against all bacteria equally, without showing specific selection of various species to any great extent. The analogy to ferment action was formulated by Buchner because of the heat sensitiveness and the instability of the bactericidal substance on standing; and he suggested that this alexin might possibly be a product of the tissue or blood cells, possibly leukocytic in origin.

Buchner found that the action of the ferment-like alexin upon bacteria was most marked at the temperature of the body, and that it was capable of destroying bacteria in the subcutaneous tissues and the serous cavities of the animal body, without the aid or coöperation of cellular elements. He inferred that there was a direct relation between the potency of the alexin and resistance against infection.

The next great step in the understanding of the bactericidal processes was now made by Pfeiffer as a consequence of studies upon the nature of cholera immunity. Pfeiffer^{11 12} found that the injection of cholera spirilla into the peritoneal cavity of a guinea pig which had recovered from a previous cholera infection was followed by a rapid destruction of the bacteria. If small quantities of exudate were taken out of the peritoneum at varying intervals after the injection, a granular change and swelling of the bacteria were noticed, followed, soon after, by complete dissolution and disappearance. Such animals would recover from doses of bacteria which, in control animals of the same weight, resulted in death. He further found that the phenomenon was specific, in that the dissolution of cholera organisms only occurred in the cholera-immune animals, other bacteria being unaffected. In other words, the guinea pig had acquired a specific antibacterial power, expressed by the process of "bacteriolysis," a property possessed to only a very slight extent by the peritoneal exudate of a normal animal. It was the next logical step to determine whether the bacteriolytic power could be transferred to the peritoneal cavity of a normal animal by injecting, together with the bacteria, a small amount of the serum of such an

¹¹ Pfeiffer. *Zeitschr. f. Hyg.*, Vol. 18, 1894; also Vols. 19 and 20.

¹² Pfeiffer & Isaëff. *Deutsche med. Woch.*, No. 18, 1894.

immune animal. This was indeed found to be the case and, although such immune serum, like normal serum, is deprived of its *in vitro* bactericidal power on heating, Pfeiffer found, in his intra-peritoneal experiments, that heated serum is quite as effectual as fresh immune serum in transferring passive immunity to a normal guinea pig. We may summarize the important harvest of facts obtained from these experiments of Pfeiffer in the following statements:

1. Rapid dissolution of cholera spirilla takes place in the peritoneal cavity of a cholera-immune guinea pig. Similar lysis takes place not at all, or only to a slight extent, in the peritoneum of a normal pig. In consequence of the lysis the immune pig will survive the injection of quantities of bacteria which invariably kill normal animals of the same weight.

2. The protection obtained in this way is specific.

3. The protection may be transferred from an immune to a normal guinea pig, by injecting a little immune serum together with the bacteria into the peritoneum of the normal animal. In a normal animal so treated lysis is in every way similar to that observed in the immune pig.

4. The transfer of the lytic power and consequent immunity can be brought about not only by means of fresh immune serum but by heated serum as well, although the latter has lost all its alexic power because of the heating.

Of the phases of this "Pfeiffer phenomenon" the one most difficult to understand, in the light of the knowledge of that time, was the transference of the lytic property with the heated serum. Pfeiffer very naturally took his experiments to signify that the actual destruction of bacteria in the animal body could take place entirely without the phagocytic participation of the body cells, a view in sharp contrast to that of the Metchnikoff school, and based upon his observation of the complete extracellular disintegration of the spirilla in the peritoneal exudate. He assumed, however, that there was an indirect participation on the part of the cells. The observation that heated serum, inactive outside the body, was efficient when introduced into the peritoneum, persuaded him that the coöperation of the living tissues was a necessary factor, and he assumed a possible activation by substances derived from the endothelial cells lining the peritoneal cavity. In the same way he explained his failure to observe actual bacterial dissolution in hang-drop preparations, even when fresh serum was used in the experiment.

It will be interesting to examine a protocol of an experiment such as those carried out in the performance of the Pfeiffer phenomenon in order to make the actual occurrences entirely clear. In such experiments the quantity of bacteria used must be chosen with some regard to the virulence and toxicity of the particular culture em-

ployed, since, as we shall see, protection of animals by bactericidal or bacteriolytic sera does not follow the law of multiple proportions as in the case of the protection against toxins by antitoxins. While the dose of bacteria chosen should be considerably above the minimal lethal dose for an animal of the weight used, it should nevertheless be remembered that the bactericidal serum does not possess antitoxic properties against the poisons liberated or produced as the bacteria undergo dissolution, and at best the protection by bacteriolysis is limited to a very definite maximum of bacteria, beyond which no further increase of serum quantity will avail. The following table will illustrate an experiment of this kind in which, in a series of guinea pigs, the bacteriolytic protective power (titre) is determined by comparative tests.¹³

PFEIFFER PHENOMENON

| Weight of guinea pig | Dose of bacteria* cholera spirilla | Amount of inactivated immune serum | Result |
|-----------------------|------------------------------------|---|---|
| (1) 215 gm. | 2 mg. | 0.1 c. c. in 1 c. c. salt solution. | Complete dissolution in less than 1 hour. Lives. |
| (2) 230 gm. | 2 mg. | 0.05 c. c. | About the same as first. |
| (3) 200 gm. | 2 mg. | 0.01 c. c. | Somewhat slower than in other two; a few unchanged spirilla after 1 hr. Final dissolution. Pig lives. |
| (4) 245 gm. | 2 mg. | 0.005 c. c. | Similar to (3) but complete dissolution in 2 hrs. Pig lives. |
| (5) 220 gm. | 2 mg. | 0.001 c. c. | After 30 min. the spirilla seem to have begun to multiply. Dies with innumerable active spirilla in peritoneum. |
| <i>Normal control</i> | | | |
| (6) 210 gm. | 2 mg. | 0.1 c. c. normal inactive rabbit serum. | Very slight lysis at the beginning. Soon rapid multiplication. Dies. |

*The bacteria may be measured for such an experiment by standard loopfuls (1 loop being equal to 2 milligrams), or by volume in emulsion with salt solution.

Pfeiffer has established a system of standardization for the measurement of sera by this technique. He speaks of one immunity unit as the smallest amount of such a serum which is capable of causing

¹³ For extensive discussion of the technique of such tests see Boehme in *Kraus u. Levaditi Handbuch*, etc., Vol. 2, p. 366. The scheme of presentation of our example is taken from that used by him. See also Pfeiffer, *Zeitschr. f. Hyg.*, Vol. 19, 1895, p. 77.

complete dissolution of 2 milligrams of culture material¹⁴ (of a standard culture) and saving the life of the animal. The unit of the serum in the preceding test would accordingly be 0.005 c. c., and the titre of the serum, expressed in Pfeiffer's language, would be 200 units to the cubic centimeter. Owing to the great variation in the virulence and toxicity of different strains of the same organism, and also because of the difficulties opposed to the visible dissolution of many bacteria, which may be killed by the serum without showing much evidence of solution, the practical application of Pfeiffer's standardization is not universally possible. In doing experiments by this technique, whatever their purpose may be, accurate adjustment of bacterial amounts and preliminary studies of virulence must be made in order that the tests may be of real value and, failing visible lysis, the death of the animals must be taken as the indicator of the titration. Comparisons of results obtained with two different cultures of the same species are consequently of value only when the minimal lethal dose of each and its toxicity have been studied before the final tests are made.

The cardinal points of Pfeiffer's phenomenon were rapidly confirmed, but his assumption that the process could take place only within the animal body was soon corrected by both Metchnikoff¹⁵ and Bordet.¹⁶ Both of these investigators succeeded in producing extracellular lysis of cholera spirilla in hang-drop preparations. The former produced the phenomenon by adding to the hang-drop preparations small quantities of extracts of leukocytes, and thus attempted to correlate Pfeiffer's observations with his own opinions regarding the importance of the leukocytes in bacterial destruction. The latter, however, subjected the phenomenon of bacteriolysis, both in vivo and in vitro, to a careful analysis and obtained results which definitely disproved the necessity of cellular intervention in this phenomenon, and furnished facts regarding the process which stand uncontradicted to the present day. Upon the basis of these our modern views of the mechanism of cytolysis in general are founded.

Bordet showed that the bacteriolytic properties of immune serum are indeed destroyed by heating to from 50° to 60° C. If, however, to such a heated immune serum there is added a small quantity of fresh normal serum, the bacteriolytic power is restored with undiminished vigor. He recognized in consequence that there were two distinct serum elements necessary for the process. Fresh normal serum by itself had very slight or no bacteriolytic power. Fresh immune serum had powerful and rapid effects. Heated immune

¹⁴ The standard "loop" used in many laboratories for the rough measurement of quantities of bacteria from agar cultures takes up approximately 2 milligrams of the material.

¹⁵ Metchnikoff. *Ann. de l'Inst. Past.*, Vol. 9, 1895.

¹⁶ Bordet. *Ann. de l'Inst. Past.*, Vol. 13, 1899.

serum had lost its power completely, but this was restored to it by the addition of the fresh normal serum. He noted, furthermore, that the specific nature of the bacteriolysis by the immune serum was unchanged after it had been inactivated by heat and reactivated subsequently by the normal serum. The inference was plain. Immunization of an animal incites the production, in the blood of this animal, of a "preventive" substance, which is moderately resistant to heat, and which is specific for the bacteria employed in the immunization. This substance cannot act upon the bacteria alone, however, but depends for its effective functionation upon the coöperation of another substance present universally in normal serum, the "bactericidal" substance, which is non-specific, corresponds to Buchner's alexin, and is apparently not increased by the process of immunization. These are the fundamental facts revealed by the early studies of Bordet, and they are stated in the present connection merely as experimental facts, without further elaboration of the later theoretical interpretation placed upon them by Bordet himself and by Ehrlich and his followers.

In the course of these studies Bordet¹⁷ had used the immune serum produced in a goat by injection of cholera spirilla. As normal serum he had used guinea-pig serum, and the latter frequently contained a few blood corpuscles. He noticed that these corpuscles were frequently clumped in the goat serum and correlated this with the similar clumping (agglutination) of cholera organisms which he had noticed in this and other sera. In his incidental observation of the phenomenon of agglutination he had concluded that the living nature of the bacteria had no importance as far as their agglutination was concerned, dead organisms being as readily agglutinated as living.

Reasoning from this similarity between blood cells and bacteria in their behavior in serum, it occurred to him that the phenomena both of agglutination and of lysis might be expressions of general biological laws, not limited to bacteria. Accordingly he injected rabbit blood into guinea pigs, and examined the serum of animals so treated for its action upon rabbit corpuscles, *in vitro*. He found that the sera of "blood-immune" animals had acquired not only increased agglutinative power against the corpuscles injected, but had also acquired specific "hemolytic" powers, that is, the property of causing a solution of hemoglobin out of the red cells. (For the process of serum hemolysis does not consist of a complete dissolution of the red corpuscles, but rather in the liberation of the hemoglobin from the cell stromata.) The latter (shadow forms) can be recovered undisintegrated by the centrifugation of hemolyzed blood. The

¹⁷ See Bordet's own account in a "Résumé of Immunity"; "Studies in Immunity," Bordet, collected and translated by Gay, Wiley & Son, N. Y., 1909.

process, like that of bacteriolysis, was specific in that the hemolytic power was lost if the serum was heated to from 50° to 60° C., but could be restored undiminished by the addition of a little fresh normal serum, in itself possessing no hemolytic properties for the given species of cell. The specificity of the phenomenon again was seen to reside entirely in the heat-stable factor, the heat-sensitive or "alexin" factor being non-specific, and not increased during the process of immunization.

Observations related to those of Bordet concerning hemolysis were made independently, in the same year, by Belfanti and Carbone, who had observed that the serum of animals treated with blood cells of another species became toxic for this species, and extensive confirmation of the phenomenon of hemolysis was obtained, in the year following, by the work of von Dungern, and by that of Landsteiner.

After Bordet had thus established the important fact that hemolysis was in every way analogous to bacteriolysis in that, like bacteriolytic sera, hemolytic sera could be inactivated by heat, but reactivated by the addition of small quantities of fresh normal serum, Ehrlich and Morgenroth¹⁸ undertook an elaborate study of the mechanism of hemolytic phenomena, hoping thereby to elucidate the mechanism of lysis in general. For it is obvious that hemolysis lends itself far more easily to experimentation than does bacteriolysis, and, as we shall see, experiments on hemolysis can be made with a considerable degree of accuracy. Ehrlich and Morgenroth approached the investigation of the hemolysins from the point of view of the side-chain theory, formulated by Ehrlich in connection with his work on the toxins. According to this theory, it will be remembered, the hemolytic substances in the sera of animals treated with blood corpuscles represent the receptors or side chains of tissue cells. These receptors were originally integral chemical elements of the body cells, by means of which the cell became united to the injected erythrocyte (or bacterial) protein. Since union with the foreign substance blocked these receptors or side chains, thereby rendering them useless, they had been regenerated and, under the influence of immunization, regenerated in excess, cast off by the cell, and were now free in the blood stream as hemolysins (or bacteriolysins).

If this conception of the process was the correct one, Ehrlich and Morgenroth argued, the hemolytic substances of any immune hemolytic serum should possess specific chemical affinity, "haptophore groups," as they expressed it, for the blood cells which had been used in the immunization.

In order to show this, they inactivated at 56° C., by the method of Bordet, a goat serum which was hemolytic for beef blood, left it

¹⁸ Ehrlich and Morgenroth. *Berl. klin. Woch.*, Nos. 1, 21, and 22, 1900.

in contact with beef blood corpuscles for 15 minutes at 40° C., and then separated the cells from the supernatant fluid by centrifugation. To the blood cells they then added a little normal goat serum (by itself not hemolytic for beef blood) and found that complete hemolysis occurred. The addition of normal goat serum and beef blood cells to the supernatant fluid, however, resulted in no change.

In the following diagram we have tried to represent this basic experiment, giving the facts only of the experiment without using any of the usual symbols which imply agreement with a theory.

EXPERIMENT TO SHOW THAT THE ANTIGEN (IN THIS CASE RED BLOOD CELLS) ABSORBS THE SPECIFIC HEAT STABLE ANTIBODY OUT OF THE IMMUNE SERUM.

In a test tube $\left\{ \begin{array}{l} 4 \text{ c. c. of } 5 \text{ per cent. emulsion of washed beef blood.} \\ 1 \text{ c. c. of inactivated blood serum of a goat treated with beef blood.} \end{array} \right.$

These substances are left together at 37.5° C. for one hour and then centrifugalized into:

I

Sediment of Corpuscles.—To this are added 4 c. c. salt solution and 0.8 c. c. fresh normal goat serum, by itself not hemolytic for beef corpuscles.
Result = *Complete hemolysis.*

II

Supernatant Fluid Containing the Serum and Salt Solution.—To this are added washed beef corpuscles and 0.8 c. c. fresh normal goat serum.
Result = *No hemolysis.*

Summarized, together with the facts we have already outlined, this basic experiment has the following significance: the fresh serum of the goat, previously injected ("immunized") with the beef blood, possessed the property of dissolving the hemoglobin out of beef corpuscles, viz., hemolyzing them. Heating this serum to 56° C. for 20 minutes, as Bordet has shown, deprives the serum of all hemolytic power, i. e., inactivates it. The addition of a little fresh goat serum, in itself inactive, completely reactivates the hemolytic properties of the heated immune serum. So far, as we have already seen, this shows that hemolysis is a dual process in which a heat-sensitive and a heat-stable substance co-operate, neither of them capable of producing lysis by itself. The heat-sensitive ingredient, corresponding to Buchner's "alexin," is present in normal serum, and, as Bordet¹⁹ and von Dungern²⁰ had shown, is not increased in the process of immunization, and is apparently not specific. The heat-stable substance, therefore specific and increased in immunization, must represent the receptors, overproduced and cast off into the circulation. And, as Ehrlich and Morgenroth have now shown in the experiment just described, this heat-stable element is actually bound to the red corpuscles, and renders them susceptible to the action of

¹⁹ Bordet. *Ann. de l'Inst. Past.*, Vol. 12, 1898.

²⁰ v. Dungern. *Münch. med. Woch.*, No. 20, 1900, p. 677.

the heat-sensitive substance in the normal goat serum. And furthermore, in attaching to this heat-stable element, the blood cells have removed it from the solution. For we have seen, in the experiment, that addition of corpuscles and normal serum to the supernatant fluid resulted in no hemolysis, showing that the third necessary element, originally in the mixture, had been carried down with the red cells.

In these and other experiments then, it was shown that only the heat stable substances could be fixed by the red cells, and this even at temperatures at or about 0° C. (a fact which indicates the strong affinity between the two substances), while the heat-sensitive "alexin," which Ehrlich now called "*complement*," could not attach directly to the red cells. For if such complement, in the form of fresh serum, was added to washed red blood cells, and the mixture after standing at 40° C. for some time was centrifugalized, the complement remained in the supernatant fluid, as could be easily shown by an experiment such as the one represented in the following protocol.

EXPERIMENT TO SHOW THAT COMPLEMENT OR ALEXIN IS NOT ABSORBED BY
UNSENSITIZED CELLS

Mixed in a test tube { 4 c. c. of 5 per cent. emulsion of washed beef blood.
0.8 c. c. of fresh normal goat serum (alexin or complement), not, by itself, hemolytic for beef blood.

These substances are left together at 37.5° C. for one hour, then centrifugalized into:

I
Sediment of Cells.—To this is added *inactivated serum* of immune goat which would cause hemolysis if alexin were present.

Result = *No hemolysis*.

II
Supernatant Fluid (salt solution and serum).—To this is added washed beef blood and inactivated serum of immune goat containing heat stable element.
Result = *Complete hemolysis*.

Although, therefore, the red cells bind the thermostable specific antibody of the immune serum and not the complement or alexin, it was shown both by Bordet and by Ehrlich and his collaborators that the red cells, after absorption of the thermostable substance, when exposed to the action of the complement, were not only disintegrated by hemolysis but, in the process, fixed or attached the complement, so that this was no longer available for further activation of other sensitized cells.

The fact that the alexin or complement is used up during processes of lysis, as first described by Bordet, Ehrlich, and others, has recently been made the subject of repeated investigation, since this is out of keeping with the general enzyme or fermentlike nature of complement indicated by many of its other properties.

Muir,²¹ who studied the conditions thoroughly, comes to the conclusion that the complement is in truth used up in hemolysis, but that it does not always disappear completely, this depending upon the relative amount of sensitizer or amboceptor present. (He confirms the quantitative ratios between the two substances found by Morgenroth and Sachs in hemolytic reactions, a subject discussed by us in another place.)

Liefmann and Cohn,²² in a more recent publication, have come to different conclusions. They believe that the disappearance of free complement from hemolytic complexes is not due to its chemical union with the sensitized cells in the process of hemolysis, but is due rather

(1) to a fixation by the products of hemolysis (stromata, etc.) after the reaction is accomplished,

(2) to dilution, and

(3) to weakening because of prolonged preservation in dilute solution at 37° C.²³

Theoretically this is of considerable importance if confirmed, since it would bear out strongly the conception of complement as a true enzyme or ferment. From the point of view of the practical utilization of complement fixation for various purposes it makes little difference, since here the disappearance of complement is the essential thing, irrespective of whether this occurs in the course of its activity or because of fixation by the products of its own action.

We now have the basic principle of hemolysis; facts which can easily be shown to hold good for bacteriolysis and for the bactericidal processes even when no actual solution takes place. Briefly reviewed, these facts are as follows: The antigen (blood cells, bacterial cells, etc.) undergoes hemolysis or bacteriolysis when acted upon by two factors, one a thermostable substance, specific and increased during immunization, the other a thermosensitive substance present in fresh serum, not increased²⁴ by immunization of the animal with the antigen and not specific. The specific thermostable substance becomes united with or fixed to the antigen regardless of the presence or absence of the thermosensitive alexin or complement, and with such avidity that the union takes place even at 0° C. The alexin or complement, however, cannot enter into relation with the antigen unless this has been rendered susceptible to it by attachment to the thermostable specific substance. When this has taken

²¹ Muir. *Lancet*, Vol. 2, 1903, p. 446.

²² Liefmann and Cohn. *Zeitsch. f. Immunitätsforsch. Or.*, Vol. 8, p. 58, 1911.

²³ In the ordinary dilution used in Wassermann tests, the unit of complement employed may deteriorate entirely within several hours at 40° C.

²⁴ Bordet. *Ann. de l'Inst. Past.*, Vol. 12, 1898. Confirmed by v. Dungen, *Münch. med. Woch.*, No. 20, 1900.

place, union with complement occurs, but only at temperatures above 0° C. (the speed and completeness of the union increasing as the temperature approaches 40° C.), and the result of the union is lysis or, in the case of bacteria not easily soluble, the bactericidal effect.

Early in their researches, Ehrlich and Morgenroth were led to speculate upon the possibility of the formation of lytic antibodies within the animal against its own tissue cells. It would be of the greatest importance to pathology, as they point out, if it could be shown that an animal could produce hemolysins, for instance, against its own blood cells. Thus, if an extensive internal hemorrhage occurred from trauma or other cause, in the course of which considerable quantities of erythrocytes are subjected to disintegration and absorption, it is at least conceivable that specific "auto-hemolysins" might appear which would lead to a chronic destruction of the red cells, with consequent anemia. This form of reasoning, as we shall see, has been extensively applied in the case of the cytotoxins for the explanation of a variety of pathological conditions. Ehrlich and Morgenroth approached the question experimentally in their further work on the hemolysins in goat blood. They found that it was comparatively easy to produce hemolysins in one goat by treatment with the erythrocytes of other goats, *isohemolysins*, as they called them.

Although, however, the blood serum of such an immunized goat was strongly hemolytic, not only for the blood cells of the goats whose blood had been injected, but also for the erythrocytes of certain other goats (though not, as we shall see, for goats in general), it was never in any case active against this goat's own cells. Moreover, while the other sensitive erythrocytes could absorb the hemolytic antibody out of the inactivated serum, the insensitive corpuscles of the goat himself seemed to possess no affinity whatever for the lysin of his own serum; mixed with the serum they failed to absorb out the hemolysin. This was in no sense, therefore, an *autolysin*.

These experiments show a remarkable individual variation between the similar tissues of animals of the same species, since Ehrlich and Morgenroth were indeed able to show that the insensibility of the goat's own corpuscles depended upon a complete absence of receptors for the isolysin. For, to explain the lack of "autolytic" action of such a serum, two possibilities could be assumed. One, as above, that the corpuscles of the goat possessed no receptors by means of which the isolysin could be "anchored" or, second, that, although such receptors were present, they were already satisfied, or saturated with the lysin in the blood stream. In the latter case it would be hard to understand why hemolysis had not taken place.

In order to completely disprove the latter possibility, Ehrlich and Morgenroth did not allow the matter to rest upon conjecture, but

resorted to an ingenious method of experimentation which yielded a further important result, namely, the discovery that the injection of antibodies into animals may give rise to "anti-antibodies." They injected inactivated hemolytic serum into goats whose corpuscles were sensitive to its action, and found that an "anti-isolysin" was formed, which, mixed with hemolysin and sensitive corpuscles, prevented hemolysis. Injection of such an isolysin into the goat from which it had been obtained, however, did not yield anti-isolysin, and it was therefore reasonable to suppose that its tissue cells possessed no suitable receptors. This failure of the production of antibodies by an animal against its own tissue cell has been spoken of by Ehrlich as "Horror Autotoxicus."

These rather involved experimental data will be shown to have a more than purely academic value when we come to speak of the problems of cytotoxin formation, and although they seem to show that auto-antibodies do not form, as a rule, exceptions to this generalization have been observed. The most notable of these is the observation of Landsteiner and Donath²⁵ made in connection with the condition of paroxysmal hemoglobinuria. It was found that in such cases, in which hemoglobinuria follows exposure to cold, the blood serum of the patient contains an "autohemolysin." If the blood of such a case is taken into oxalate or citrate solution, and allowed to stand at ordinary or incubator temperature, nothing occurs. If, however, such blood is cooled to 0° to 10° C. and then warmed gradually to the temperature of the body, rapid hemolysis occurs. In this case the "amboceptor" of the serum is apparently fixed or anchored by the blood cells only at a low temperature, the complement becoming active as the blood is warmed. Although Landsteiner's observations are undoubtedly accurate, it is likely that this mechanism does not explain all such cases. The writer has had occasion to examine carefully a number of clinically diagnosed cases of this sort with a partially successful "Landsteiner" phenomenon in one of them only. Other observers have, however, confirmed Landsteiner's observation in well-established cases of the condition.

Before we leave the subject of iso-antibodies it will be interesting to discuss for a moment the existence of isolysins in animals other than goats and more especially those occurring in human beings, phenomena which have recently assumed considerable importance in view of the frequent therapeutic performance of blood transfusion.

The peculiar facts unearthed by Ehrlich and Morgenroth²⁶ indicated specific differences between red blood cells of individuals in the same species (goats), which could only be recognized by the

²⁵ Landsteiner and Donath. *Munch. med. Woch.*, 1904, p. 1590.

²⁶ Ehrlich and Morgenroth. "Über Hämolsine," *Berl. kl. Woch.*, 1900, No. 21.

development of immune-isolysins. Work on other species of animals has indicated that this fact has a broad significance and that similar differences between individuals of the same species occur in many, if not all, species of animals. Isolysins similar in principle to those of Ehrlich and Morgenroth were produced by Ascoli²⁷ in rabbits; by Todd and White,²⁸ in oxen; by Ottenberg, Kaliski, and Friedmann²⁹ in dogs; by Ottenberg and Thalhimer³⁰ in cats, and by Hada and Rosenthal³¹ in chickens. In all these instances the isolysins developed showed the same peculiarities, namely, that they attacked the cells of certain individuals and left the cells of other individuals of the same species unharmed. Recent work on the isolysins occurring naturally in the human blood has thrown considerable light on the nature of immune isolysins.

The occurrence of isolysins in human blood was first noted by Maragliano³² in 1892, and a large amount of work had been done before it was clear that the occurrence of isolysins is not a characteristic of disease. The work of Moss³³ and of Grafe and Graham³⁴ has shown that the occurrence of isolysins is parallel with that of iso-agglutinins (see chapter on agglutination), and that there are in human bloods two isohemolysinogens, A and B (corresponding to the two agglutinogens, A and B), and two isohemolysins, α and β . The hemolysinogens occur regularly according to the same rule as the agglutinogens, but the hemolysins, while they always follow the same rule when present, may be present or latent. Thus a person whose red cells contain A may or may not have β , and never has α ; a person whose red cells contain B may or may not have α , but can never have β ; a person whose red cells are susceptible to both α and β never has any hemolysin in the serum. It seems likely that the substances A and B, which cause the susceptibility of red cells to the corresponding hemolysins, are definite biochemical structures which possibly may be inherited in a similar way to the iso-agglutinogens and that similar substances (probably a larger number of them) are present in the blood cells of various species of lower animals. This readily explains the apparent irregularity attending the development of isolysins in the lower animals. The reason for the natural occurrence of such isolysins in human sera and occasionally in the sera of lower animals, however, is a complete mystery. From

²⁷ Ascoli. *Münch. med. Woch.*, 1901.

²⁸ Todd and White. *Nature*, June 23, 1910.

²⁹ Ottenberg, Kaliski, and Friedmann. *Jour. Med. Res.*, Vol. 28, 1913.

³⁰ Unpublished personal communication.

³¹ Hada and Rosenthal. *Zts. f. Imm.*, 1913, 16, p. 524.

³² Maragliano. "IX Congr. f. Innere Med.," 1892.

³³ Moss. *Johns Hop. Hosp. Med. Bull.*, March, 1910.

³⁴ Grafe and Graham. *Münch. med. Woch.*, 1911, pp. 2257, 2338.

the work of Matsuo³⁵ it seems likely that the autolysins of paroxysmal hemoglobinuria are not identical with the isolysins α and β .

Since the reintroduction of blood transfusion as a therapeutic measure the occurrence of hemolysis between the blood of two human beings has become of great practical importance. A number of serious or fatal accidents following the transfusion of hemolytic blood have been reported. Ottenberg and Kaliski³⁶ have shown that it is possible regularly to avoid such accidents by preliminary blood tests.

These tests are easily carried out by obtaining serum and washed blood cells from both prospective recipient and donor, and testing them one against the other for hemagglutination and hemolysis, as follows:

1. Active Serum Donor, 0.5 c. c. + Red Cells Recipient, 0.5 c. c.
(5% emulsion in NaCl)
2. Active Serum Recipient 0.5 c. c. + Red Cells Donor 0.5 c. c.
3. } Controls of both varieties of cells in salt solution.
4. }

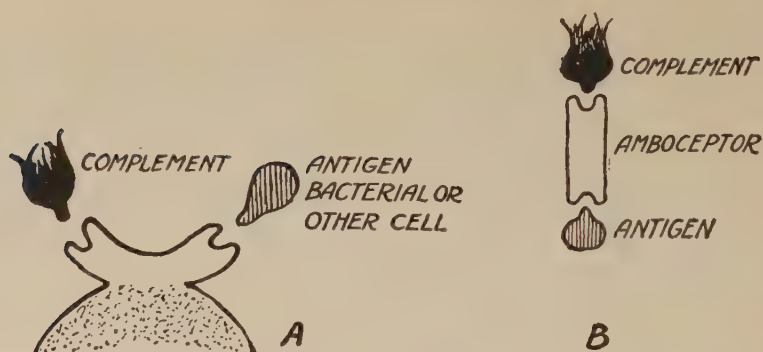
Such tests should be observed for at least two hours before final readings are taken.

Although we have by no means covered in detail the entire experimental plan followed by Ehrlich and his collaborators during their early work, we are now ready to consider the basic views on the structure of the lytic antibodies which they deduced.

It appears from the preceding that the thermostable hemolytic antibody must, of necessity, unite with the red cell before the complement or alexin can exert its action upon it. Ehrlich conceives this process as a mediation on the part of the heat-stable substance between the antigen and the alexin or complement. The heat-stable body, which he calls "amboceptor," because of its assumed mode of action, possesses two combining groups—one the "cytophile," by means of which it is anchored to the sensitive cell, the other the "complementophile," by means of which it exerts affinity for the complement. The original cell receptor, from which such an "amboceptor" takes its origin, is one which not only can combine with the antigenic substance offered for assimilation, but which also possesses another atom group by means of which it can enlist the aid of the digestive ferment of the blood, the alexin or complement. Cast off into the blood stream, as a result of overregeneration, it now appears as a "double" receptor, which can form a link between antigen and complement.

³⁵ Matsuo. *D. Arc. f. kl. Med.*, Bd. 107 H₄, p. 335.

³⁶ Ottenberg and Kaliski. *J. A. M. A.*, Vol. 61, 1913.



SCHEMATIC REPRESENTATIONS OF A RECEPTOR OF THE THIRD ORDER.

Ehrlich's conception of the relationship of antigen, amboceptor, and complement in the bactericidal and hemolytic process. In A the receptor is still a part of the body cell, in B it has been overproduced, and is free in the circulating blood.

In his general scheme of diagrammatic representation of these processes Ehrlich refers to the "amboceptors" as "haptines" of the third order.

Now it is quite plain, from the extreme specificity which results when an animal is immunized with any given variety of blood cells or bacteria, that there must be as great a variety of such amboceptors as there are different antigens, and indeed an animal immunized with two or more antigens may simultaneously contain in its blood serum a corresponding number of different amboceptors.

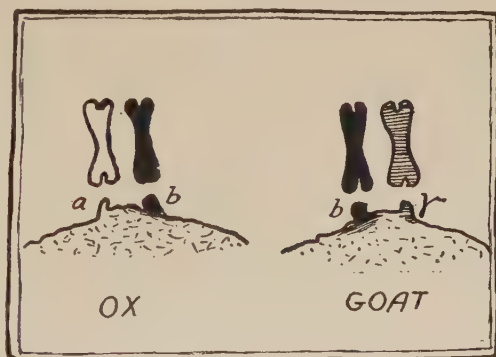
This assumption of the multiplicity of "amboceptors" in the same serum is, of course, forced upon us by the fact of specificity, and the frequently repeated observation that the same serum may contain heat-stable lytic antibodies against a variety of antigens, each antigen absorbing out of such a serum that antibody only which specifically reacts with it. This fact has, of course, never been denied, and it is a frequent misunderstanding of the views of Bordet, which will be discussed directly, to assume that he has combated the "multiplicity of amboceptor" in the sense just outlined. Ehrlich and Morgenroth, however, have expressed themselves in favor of the conception of a multiplicity of "amboceptor" not only in this sense, but as occurring in response to immunization with one and the same antigen.

Ehrlich and Morgenroth³⁷ assume that any cellular antigen, blood or bacterial cell, substances of great complexity of chemical structure, must necessarily be possessed of a large number of different side chains or receptors. When immunization is practiced with such cells a correspondingly varying number of different amboceptors must result. They found, for instance, that when rabbits are

³⁷ Ehrlich and Morgenroth. *Berl. klin. Woch.*, Nos. 21 and 22, 1901.

immunized with ox blood, the resulting antiserum is capable of producing hemolysis not only of ox blood but of goat's blood as well, though to a lesser degree. They conclude from this that the hemolytic action of the serum must be referred to the presence of at least two kinds of amboceptor, especially since repeated experiments with different anti-ox-blood sera showed that there was no regularity in the proportions of hemolysins for ox and goat blood, respectively. This opinion they further fortify by showing that exposure of the serum to ox blood deprives it of all its hemolysins, both those for ox and those for goat's blood, whereas absorption with goat's blood alone removes the specific goat's blood hemolysins only. They translate their understanding of the conditions to graphic form by the following diagram:³⁸

If ox blood is injected, α and β receptors being present, α and β amboceptors are formed, and ox blood can consequently anchor both amboceptors. The presence of β receptors in goat's blood also explains the moderate hemolysis of this blood by the antiserum, but lacking the α receptors which, in this case, represent the larger proportion, these blood cells cannot remove all the amboceptor for ox blood out of the serum.



SCHEMATIC REPRESENTATION OF EHRLICH AND MORGENROTH'S CONCEPTION OF THE COMPLEX STRUCTURE OF AN ANTIGEN.

(After Ehrlich and Morgenroth. *Berl. klin. Woch.*, Vol. 38, 1901.)

The example given, of course, represents the simplest assumed case, and Ehrlich and Morgenroth believe that the same blood or bacterial cells may possess an entire series of such receptors, some of them being dominant for the given species, others being merely secondary or "partial," in varying proportions.

If we grant the fundamental premises of Ehrlich respecting the "double receptor" or "amboceptor" nature of the specific antibody and its mediation between antigen and complement by means of a cytophile and a complementophile receptor, certain logical consequences of this conception suggest themselves, which, in their many ramifications, have been the subject of much investigation. And although many phases of these researches are no longer commonly accepted, some, indeed, being untenable in the light of more recent

³⁸ Ehrlich. "Gesammelte Arbeiten," p. 147.

discoveries, the influence of this work upon the development of immunology has been so important that it must be briefly reviewed in order that controversial questions may be justly considered.

The comparison of the action of hemolytic sera with that of ferments, and the possibility of producing antiferments by the injection of the ferments into animals, obviously suggests a similar induction of antihemolysins by the treatment of animals with lysins. This, we have seen, was the method employed by Ehrlich and Morgenroth in their studies of the causes of the failure of autolysin formation in goats. They extended this work with the purpose of ascertaining whether or not there were differences in the structure of the cytophile groups of the various amboceptors formed when various animals were injected with any given species of red blood cells. After obtaining a strong hemolytic serum by injecting ox blood into a rabbit, they treated a goat with the inactivated serum of this rabbit. The result was that the serum of the goat so treated, when mixed with ox blood cells and the hemolytic serum, prevented the sensitization of the cells by the hemolysin. They then measured the neutralizing power of such an "anti-amboceptor" or "anti-sensitizer" against a variety of hemolytic sera produced with ox blood in different animals and found that, while this "anti-amboceptor" neutralized the hemolytic action of an antiserum produced in rabbits, it had but an indifferent or entirely ineffective neutralizing power upon similar ox blood hemolysins derived from goats, geese, dogs, rats, or guinea pigs. They concluded from this that, although these various lysins had been produced in the different animals by the injection of the same antigen, viz., ox blood, and possessed affinity for the ox blood in consequence, they must necessarily differ from each other in some way, since they were not equally neutralized by the same antilysin. It seemed to them that the difference in such cases must depend upon variations in the structure of the cytophile group of the amboceptor, a conclusion which they based upon the foregoing experiments and sought to support by the following reasoning: When an animal is treated with sensitizers or amboceptors, they reasoned, these bodies react with the tissue cells by means of the cell-receptors. These receptors are then overproduced and extended into the circulation as free atom-groups.

They now act as "anti-amboceptor," free in the serum, but are in structure merely overproduced cell receptors, identical with those which originally united *on* the cell with the injected amboceptor.

Ehrlich and Morgenroth,³⁹ therefore, believed that the neutralization of the amboceptor by the antilysin depended upon a union of the latter with the "cytophile" group of the former, preventing its subsequent union with the red cells. And since one and the same antilysin did not thus invalidate the action of all the amboceptors

³⁹ Ehrlich and Morgenroth. *Berl. kl. Woch.*, No. 22, 1901, p. 600.

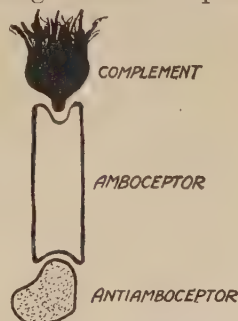
for ox blood (derived from different animals), they concluded that these "amboceptor" must possess different "cytophile groups."

That this conclusion of Ehrlich and Morgenroth is not correct seems to follow the subsequent work of Bordet.⁴⁰ He demonstrated that it is not necessary to inject animals with specific hemolytic sera in order to obtain antilytic sera, but that the same object may be attained by injecting animals with the normal serum of an untreated animal. Moreover, if an "antisensitizing" serum so produced was added to corpuscles which had already absorbed "amboceptor," it prevented the subsequent union of these sensitized cells with alexin or complement. From this it becomes clear that, in the first place, the antisensitizer or anti-amboceptor cannot be identical with the cell receptors of the corpuscles, and, further, that the inhibition of the hemolysis which such an antisensitizer exerts, cannot be due to union with the "cytophile" group. This both contradicts the Ehrlich conception of the mechanism of "anti-amboceptors" and invalidates his argument, in this instance, in favor of the plurality of the amboceptors produced by the injection.

Bordet's experiments were later confirmed by Ehrlich and Sachs,⁴¹ who admit the error of the former "anticytophile" interpretation of Ehrlich and Morgenroth's experiments, but they still maintain that Bordet's experiments do not disprove the conception of an "amboceptor" or "Zwischenkörper" of Ehrlich. They claim that Bordet's results merely prove that the anti-amboceptor or anti-sensitizer is "anticomplementophile" instead of "anticytophile."

The principles involved we will discuss in another place in connection with Moreschi's analysis of the "anticomplements." However this may be, we may conclude that Ehrlich and Morgenroth's differentiation of amboceptors or sensitizers by the cytophile group is no longer valid.

The studies of Bordet on the antisensitizers (anti-amboceptor) had important results apart from their refutation of Ehrlich and Morgenroth's opinion. In addition to showing that such antisensitizer did not represent cell receptors identical with those that anchored the sensitizer (amboceptor) to the red blood cells, his experiments revealed the fact that such an antisensitizer neutralizes un-



SCHEMATIC REPRESENTATION OF EHRLICH AND MORGENROTH'S CONCEPTION OF THE NEUTRALIZATION OF A HEMOLYTIC SERUM BY ANTILYSIN OR ANTIAMBOCEPTOR, REACTING WITH THE CYTOPHILE GROUP. (Ehrlich and Morgenroth, *loc. cit.*)

This conception, as we shall see, has become untenable.

⁴⁰ Bordet. *Ann. de l'Inst. Past.*, Vol. 18, 1904, p. 593.

⁴¹ Ehrlich and Sachs. *Berl. klin. Woch.*, No. 19, 1905.

specifically various specific sensitizers as well as normal antibodies in the serum of the same animal; and this showed that there is no necessity of assuming a variety of specific antisensitizers, as had been done by the German workers.

As regards the multiplicity of amboceptor or sensitizer, however, though the proof of this, by means of anti-amboceptors, has had to be abandoned, as we have seen, there is still a great deal of evidence advanced in favor of such an assumption. The chief support for such an opinion is found in the "group reactions" among bacteria, similar to those observed for blood cells by Ehrlich and Morgenroth, and described above (see page 151). For it is frequently observed that the antibodies produced by immunization with one species of bacteria may have a certain though lesser degree of action upon other related forms, these in turn absorbing only a part of the amboceptor out of the serum, while the species originally used for immunization takes out all the amboceptor present. Considering the great chemical complexity of the bacterial or tissue cells, moreover, we may well expect such multiplicity. And it is, indeed, entirely reasonable to suppose that a structure as complex as the bacterial cell may contain a number of antigens and consequently give rise to a number of sensitizers which differ in that each is specific for its particular antigen only. This is merely a restatement of the phenomenon of specificity and has, as a matter of fact, no modifying influence on the general principles involved.

From the point of view of a general understanding of the processes of immunity, however, the question of multiplicity of sensitizer is not so fundamentally important as is the similar controversy which has been waged regarding the unity or multiplicity of alexin or complement. Here again there has been some misconception as to the meaning of those who maintain the unity of alexin. Neither Bordet, nor anyone else familiar with experimental conditions, has ever maintained that the alexins of different animals were functionally identical. It is a well-known fact that the fresh blood sera of various animal species differ from each other considerably in their power to activate bactericidal or hemolytic systems. In regard to hemolysis, fresh guinea-pig serum is very powerful in activating many sensitized blood-cell complexes, but weak in activating sensitized guinea-pig corpuscles. Often one finds that the alexin of an animal is entirely impotent or but weakly capable of producing hemolysis of the sensitized cells of its own species, though this is not a general rule.

Again, even without such species relationship, a given alexin may be very weak for certain complexes and strong for others. The alexin of horse blood can even be fixed to sensitized cells⁴² without

⁴² For the sake of clearness it may be repeated here that by sensitized cells we mean cells which have absorbed specific "amboceptor" or "sensitizer," and have thereby become amenable to the action of alexin or complement.

producing much, if any, hemolysis.⁴³ An alexin which may be strong for a given hemolytic complex may be weak for certain bactericidal complexes, or *vice versa*. Thus there is a large mass of evidence which shows that no two alexins are exactly alike, though the difference between them can, of course, be defined functionally only.

The difference between the opinions of Ehrlich and his school on the one hand, and the followers of Bordet, on the other, revolves not about this point, upon which all agree, but about the question of whether one and the same serum may contain more than one alexin or complement. Ehrlich and Morgenroth⁴⁴ and Ehrlich and Sachs⁴⁵ have brought forward evidence from which they deduce the existence of a number of different alexins or complements for hemolytic complexes in the same serum. The earlier experiments of Ehrlich and Morgenroth on this question were carried out by means of the filtration of normal goat serum through Pukall filters;⁴⁶ in these it appeared that the serum which passed through the filters was complementary for sensitized guinea-pig cells, while that part which had, in the original serum, activated sensitized rabbit cells was left behind. Similar differentiation of complement they later based upon experiments with anticomplementary sera which, they showed, did not equally neutralize all the complementary functions of a serum.

In support of their contention Neisser⁴⁷ described two complementary substances in rabbit serum, the one active for bactericidal complexes, the other for hemolytic, and similar experimental evidence has been brought forward by Wassermann⁴⁸ for guinea-pig and by Wechsberg⁴⁹ for goat serum.

The evidence advanced by these writers is based chiefly on experiments in which it was found that a normal serum which possessed both bactericidal and hemolytic powers could be deprived of the complement for one or the other of these activities only, by absorption with the respective cells. In addition to this, Ehrlich and Morgenroth, Ehrlich and Sachs,⁵⁰ Wendelstadt,⁵¹ and others, claimed to have differentiated various complements in the same serum by careful heating, by the action of weak acids or alkalis, or such methods as the digestion of sera by papain.

⁴³ Browning. *Wien. klin. Woch.*, No. 15, 1906.

⁴⁴ Ehrlich and Morgenroth. *Berl. kl. Woch.*, No. 31, 1900.

⁴⁵ Ehrlich and Sachs. *Berl. kl. Woch.*, No. 21, 1902.

⁴⁶ Sachs. *Berl. kl. Woch.*, Nos. 9 and 10, 1902.

⁴⁷ Neisser. *Deutsche med. Woch.*, 1900, p. 790.

⁴⁸ Wassermann. *Zeitschr. f. Hyg.*, 37, 1901.

⁴⁹ Wechsberg. *Zeitschr. f. Hyg.*, Vol. 39, 1902.

⁵⁰ Ehrlich and Sachs. *Berl. kl. Woch.*, Nos. 14 and 15, 1902.

⁵¹ Wendelstadt. *Centralbl. f. Bakt.*, I, Vol. 31, 1902.

As a rule, these experiments have been carried out with normally hemolytic serum and unsensitized cells, though in certain cases Ehrlich has employed sensitized cells; but whenever this was done exposure to complement for purposes of absorption has been for much briefer periods than when normal serum was used. This point is significant when we come to consider the objections to the interpretation of the preceding experiment in favor of a plurality of complement, objections raised chiefly by Wilde⁵² and by Bordet.

Wilde refuted particularly the experiments of Neisser, who claimed that the absorption of fresh rabbit serum with anthrax bacilli deprived this serum only of its bactericidal but not of its hemolytic complement. Wilde showed that, if a sufficient excess of anthrax bacilli (or in given cases of typhoid bacilli or cholera spirilla) were added, both bactericidal and hemolytic complement could be absorbed from normal serum. He concludes that there is actually only one alexin present, but that the red cells and anthrax bacilli differ in their susceptibility to this alexin (or, in other words, that the sensitization of these cells by the normal serum is unequal, a conclusion which seems rational in view of the fact, now well known, that one and the same complement may differ greatly in the degree of its activity upon different sensitized complexes.

Bordet has analyzed the conditions in a similar way. He found that absorption of normal serum with unsensitized cells rarely deprived this serum of all of its alexin, even when these cells were used in considerable amounts. This he attributed to the feeble sensitization of the cells. If, however, strongly sensitized cells were added to such a normal serum, all the alexin would be taken up. He refers the phenomenon of specific alexin absorption, observed by previous workers, to insufficiency in the perfection of sensitization on the part of the cells used in the preliminary exposure; and subsequent work with complement fixation seems to bear him out.

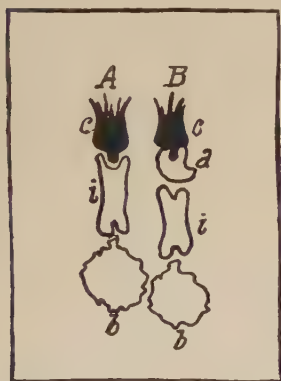
Most of these arguments, though they seem to us perfectly valid in the light of the experimental facts, have been answered by Ehrlich and his school by the assumption of the existence of so-called "poly-receptors." Ehrlich now admits that the amboceptors cannot be shown to differ from each other. However, he does not believe that differences in the intensity of sensitization explain variation in the functional efficiency of different complements upon sensitized cell complexes, nor does he accept, for proof of this, the fact that complement may be entirely absorbed out of a serum by a complex, even though the complement may be comparatively inefficient as an activator in the given case. He assumes that the sensitizer or "amboceptor" may possess a number of complementophile groups (poly-receptors), by means of which a number of different complements may

⁵² Wilde. *Habilitations Schrift*, Munich, 1901. Also *Berl. kl. Woch.*, No. 34, 1901.

become active in the given case. Thus, although such a polyeceptor, of course, is capable of uniting with the complement which activates the dominant complement, it is capable also of union with a number of other complements which have slight or no functional action whatever—the non-dominant complements. This opinion is rendered diagrammatic by Ehrlich and Marshall⁵³ in the following way:

If one carefully considers the reasons advanced for the assumption of the existence of such polyeceptors it does not seem that they are sufficiently forcible to lead one to desert the much simpler explanation of Bordet.

Related to the problems discussed in connection with the production of “anti-amboceptors” or “anti-sensitizers” are those which have arisen regarding the existence of “anti-complement” or “anti-alexins.”

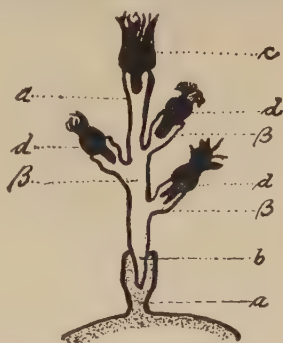


EHRLICH AND MORGENROTH'S CONCEPTION OF THE ACTION OF ANTI-COMPLEMENT.

- A. Scheme of Hemolysis.
B. Action of Anticomplement upon Hemolysin.
b. = blood cell, c. = complement, i. = immune body, a. = anticomplement.

The complementoids are not included in the scheme, since in this case they are without influence.

with the complement. There was apparently no union of the protective substance with the “complementophile” group of the ambo-



POLYECEPTOR ACCORDING TO EHRLICH AND MARSHALL.

- (a) Receptor of the Cell.
(b) Haptophore Group of the Amboceptor.
(c) Dominant Complement.
(d) Secondary Complements.
(e) Complementophile Groups of the Amboceptor:
(1) for the Dominant Complement.
(2) for the Secondary Complement.
(After Ehrlich and Marshall, *Berl. klin. Woch.*, No. 25, 1902.)

Ehrlich and Morgenroth claimed that, by the injection of active horse serum into a goat, they had obtained substances in the goat serum which neutralized horse complement. They believed that the “anti-complements” thus produced neutralized the complement by uniting with its haptophore group, thus preventing its combination with the “complementophile group” of the amboceptor. This was their conclusion because they found that the “anti-complementary” serum exerted no protective influence upon sensitized cells, when these were exposed to the serum and then removed, but that it protected against hemolysis when added to the cells together

⁵³ Ehrlich and Marshall. *Berl. kl. Woch.*, No. 25, 1902.

ceptor, but the protecting substance *did* act in direct antagonism to the complement itself.

From the fact that similar anticomplements could be produced when inactivated serum was injected into animals, they concluded that, on inactivation, there was not a complete destruction of the complement, but that during the process of heating the zymophore group of the complement only was injured, the "haptophore group," by means of which union to the tissue elements would take place, and through which, therefore, specific antibody production would be incited, remaining intact. Such altered complement they speak of as "complementoid."

Bordet has made similar observations upon the production of anti-alexins by the injection into animals both of active and of inactive serum, but in the light of further researches, which will be discussed in connection with the problems of alexin-fixation, chiefly those of Moreschi and of Gay, we are forced to the conclusion that the existence of true anticomplements is by no means certain, and that the older evidence in their favor is found to be unconvincing at the present time.

In the preceding paragraphs we have emphasized the conceptions of the cytolytic phenomena formulated by Ehrlich and his followers, and although we have brought out, whenever possible, the objections of other investigators to many of these opinions, we have not yet followed out in a systematic manner the reasoning of any of Ehrlich's opponents. In opposition to the views of his school the leading position has been taken by Bordet, who, after all, furnished in his investigations the fundamental facts which have led to a comprehension of the cytolytic processes. In explaining Bordet's views we can do no better than to follow out his own exposition as set forth in his article, "A General Résumé of Immunity,"⁵⁴ published with a collection of his papers. He expresses himself, in substance, as follows:

That the antigen, in the form of bacteria, blood cells, or cells of any other nature, meets in the body of the treated animal a "receptor" complex with which it unites is, of course, plain and agreed to by everyone. That the antibody produced by the tissues in response to such union of antigen with receptor is a direct product of the cells containing the receptors is likely. It is by no means certain, however, or, at any rate, it has never been experimentally demonstrated, that, as Ehrlich maintains, the antibody is identical with the original receptor by which the antigen was fixed or anchored to the tissue cell. It might be assumed with equal justice that the cells of the immunized animal could build up a new substance, not identical with the receptors, in consequence of stimulation by the antigen. It is also by no means certain whether the injected antigen reacts with the body cells themselves or with the normal antibodies which we

⁵⁴ "Studies in Immunity" by Bordet and collaborators. Gay, Wiley & Sons, N. Y., 1909.

know to exist in many cases. Thus the blood serum of goats may normally often contain hemolysins against rabbit corpuscles. Is it not reasonable to suppose that possibly these may furnish the point of attachment and the source of further antibody production when rabbit cells are injected into goats? In criticism of Ehrlich's assumption of the mode of action of heat-stable lytic antibody, Bordet very justly maintains that no proof whatever exists of the "amboceptor" nature of this substance. All that is certain is that the stable substance must unite with the antigen before the alexin or complement can exert its action upon it or be fixed by it. There is no entirely valid proof of the existence in this antibody of a "complementophile" and a "cytophile" group, and no satisfactory instance has been observed in which alexin has united with a heat-stable antibody which has not previously been united with an antigen.⁵⁵ All that has been shown is that the antigen, together with its specific antibody, forms a complex which has an avidity for alexin, a complex which is "endowed with properties of absorption for complement which neither of its constituents alone possesses."



SCHEMATIC REPRESENTATION OF BORDET'S VIEW CONCERNING THE INABILITY OF COMPLEMENT TO UNITE WITH EITHER ANTIGEN OR SENSITIZER ALONE AND ITS ABILITY TO BE FIXED BY THE COMPLEX FORMED WHEN THE ANTIGEN IS SENSITIZED.

Compare this figure with that representing Ehrlich's conception of the same process.

Bordet speaks of the "amboceptors," therefore, as "sensitizers," meaning by this that the antigen, by union with its antibody, is sensitized to the action of the alexin. The term "sensitizers" in no way, therefore, implies a preconceived notion, experimentally unproved, of the mode of action or structure of the sensitizer. Since we have graphically explained Ehrlich's opinions, a similar diagrammatic representation may be permitted of Bordet's opinion of the same process of union of antigen and heat-stable antibody with the consequent development of alexin-fixing property.

In this diagram the ability to absorb or unite with complement becomes evident only after a complex has been formed by the union of the two elements, antigen and antibody. The diagram must not be assumed to mean that the notch into which the complement fits symbolized necessarily an "atom group," but merely expresses the idea of "ability to absorb alexin," not assuming that this ability is

⁵⁵ Refer also to the discussion of the conglutinins at the end of this chapter.

either chemical affinity by means of a definite atom group or a mere physical change of molecular equilibrium permitting a specific complement absorption.⁵⁶

It will be seen from the preceding that the controversy between Ehrlich's "amboceptor" conception and the "sensitization" idea of Bordet turns largely upon the existence of a so-called complementophile group of the thermostable antibody. For if it were the case that this antibody possessed an atom group which permitted it to unite with alexin, independent of previous union with antigen, it would go far to support Ehrlich's view. One of the strongest arguments brought into the field in favor of such an occurrence by Ehrlich's followers is the phenomenon of Neisser and Wechsberg, which is usually spoken of as "complement deviation" (Komplement Ablenkung).

In order to make the conditions underlying this phenomenon clear, it will be of advantage to consider for a moment the methods of determining quantitatively the amount of bactericidal antibody (sensitizer amboceptor) in any given immune serum, since it was in working with such titrations that Neisser and Wechsberg made their observations.

In carrying out such measurements, it is customary to add in series, to constant amounts of bacteria, varying amounts of inactivated antiserum and constant amounts of complement or alexin. These mixtures are set away in the thermostat for 3 to 4 hours, are then mixed with agar and plates are poured. The colonies which develop will give an indication of the number of bacteria killed in each mixture when compared with similar plates poured from tubes in which the same original amounts of bacteria had been mixed with alexin alone. The following table will exemplify such a test:

| Typhoid bacilli | Typhoid antiserum inactive | Alexin | Result in colonies after 3 hours, incubation |
|---------------------------------|----------------------------|-----------|--|
| Constant quantity..... | .1 c. c. | .07 c. c. | Many thousand |
| Constant quantity..... | .01 c. c. | .07 c. c. | Many thousand |
| Constant quantity..... | .005 c. c. | .07 c. c. | 150 colonies |
| Constant quantity..... | .001 c. c. | .07 c. c. | 200 colonies |
| Constant quantity..... | .0005 c. c. | .07 c. c. | 800 colonies |
| Control I, constant quantity.. | | .07 c. c. | Many thousand |
| Control II, constant quantity.. | | | Many thousand |

⁵⁶ The diagram on page 159, though possibly not expressing with absolute accuracy the idea of sensitization, was devised because it will remove what seem to the writer frequent misconceptions of Bordet's views. Statements are found in the literature which imply (Ehrlich, "Kraus und Levaditi Handbuch," Vol. 1, p. 8) that Bordet assumes "dass das Komplement direkt an die Zelle angreift," and deny that there is experimental evidence to support this. It is perfectly true that there is no evidence to show such "direktes

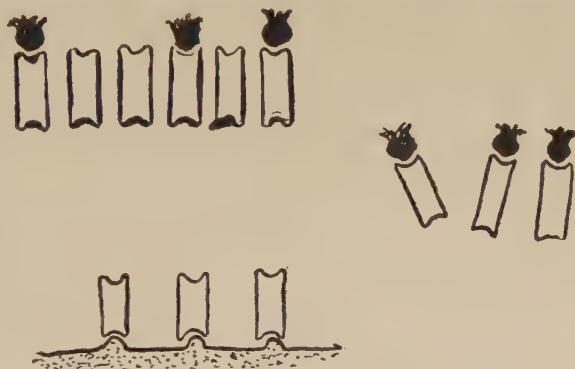
In this table it is noticeable that, although there has been considerable bactericidal action in the mixtures in which 0.005, 0.001, and 0.0005 c. c. of antiserum were used, the mixtures in which as much as 0.1 and 0.01 c. c. were present, and in which one would naturally expect a still greater antibacterial action, the contrary occurred. This surprising and curious phenomenon, showing that an excess of

antibody could actually be harmful to the functioning of the bactericidal complex, was explained by Neisser and Wechsberg by the following reasoning. In tests like the one given above a limited amount of bacteria and alexin has been mixed with the enormous amount of antibody represented in the immune serum. Although

bacteria can absorb more of this antibody than is necessary for their solution or destruction, nevertheless the higher concentration given in the table will contain quantities of "amboceptor" so far in excess of the amount that can be absorbed that much of it must remain free in the fluid. Now this amboceptor, possessing a complementophile group, is able to anchor complement or alexin as well as that which has become united with the bacteria. In consequence, there being only a limited amount of complement, some of this is deviated from the amboceptor-antigen complexes by the free amboceptor, and is, in consequence, ineffective so far as bactericidal action is concerned. In the higher dilutions of the antiserum, in which no such excess is present, the complement will be concentrated upon the "attached" or "anchored" amboceptor, and greater efficiency will result. Graphically Neisser and Wechsberg express their idea in the figure which we reproduce.

As to the accuracy of the observations of Neisser and Wechsberg there can be no question, and everyone who has occasion to carry out

"Angreifen" upon the unaltered cell, but there is evidence that this union takes place after the cell has absorbed the antibody, and no satisfactory evidence to show that the thermostable body is an intermediary, that is, forms a link as conceived in the amboceptor idea.



COMPLEMENT DEVIATION AS CONCEIVED BY NEISSER AND WECHSBERG.

The complement being united to the unbound amboceptor is thereby deviated from the amboceptor, which has gone into relation with the antigen.

(After Neisser and Wechsberg, *Munch. med. Woch.*, 1901, p. 697.)

bactericidal tests with any frequency is sure to meet with the phenomenon again and again. But their explanation, which involves the assumption of union between free sensitizer or amboceptor, and alexin or complement, without the participation of antigen, cannot be accepted since, search as we may, through the extensive experimentation that this problem has inspired, there is no instance on record in which indisputable evidence of such an occurrence has been advanced. On the contrary, there is a mass of satisfactory evidence available which indicates clearly that amboceptor or sensitizer alone cannot absorb alexin, and the Neisser-Wechsberg explanation seems consequently to be merely an interesting and cleverly conceived but improbable possibility.

What, then, is the explanation of the diminution of bactericidal effect in the presence of an excess of sensitizer? We will see that, in the study of agglutinin and precipitin reactions, phenomena exactly analogous to the Neisser-Wechsberg effect have been noticed, in the case of the agglutinins, the so-called "pro-agglutinoid" zone being a case in point. For these phenomena, as well as for that of Neisser and Wechsberg, explanations have been advanced by the Ehrlich school, similar in principle in that they all depend upon more or less arbitrary assumptions regarding affinity between the reacting bodies. Such explanations, though not outside the realm of possibility, have, however, lost much force since it has been recognized that the reactions between serum antibodies and their antigens, in general, take place according to laws far more closely analogous to those governing reactions between colloids than to those governing chemical reactions in which the laws of definite proportions can be applied. And, indeed, the reacting substances in antigen-antibody complexes are, beyond doubt, of the nature of colloids. Now, in many precipitations resulting when two colloids are mixed, an excess of one or the other factor will completely inhibit the occurrence of the precipitation; the reaction taking place only when definite proportions between the reacting bodies are present. The occurrence of such inhibition zones, due to an excessive concentration of one reagent, can be shown for agglutination and precipitation, exactly as it can in ordinary colloidal reactions, and it is more than likely that the Neisser-Wechsberg phenomenon is merely an example of a similar phenomenon.

Looked at from this point of view, far from supporting the supposition of a separate complementophile group and therefore of the "amboceptor" nature of the heat-stable lytic antibody, the Neisser-Wechsberg phenomenon indeed becomes rather a strong argument in favor of Bordet's views, and against those of Ehrlich. For, by introducing the analogy between the lytic and bactericidal processes with colloidal reactions, it takes away much force from the supposition that antigen-sensitizer alexin reactions take place according to

laws of definite proportion, an idea which still underlies, though somewhat loosely, many of the more important views of antigen-antibody reactions as conceived on the basis of the amboceptor theory.

Gay has suggested also that the Neisser-Wechsberg phenomenon may well be explicable on the basis of the fixation of complement by precipitates. In a succeeding section we will discuss the fixation of alexin, which occurs when a dissolved protein is brought together with its specific antiserum. It is not impossible that this may occur when bacterial emulsions, from which a small amount of bacterial protein may well go into solution, are brought together with antiserum in concentration. Under such conditions a reaction might readily occur which would lead to the fixation of alexin and its consequent deviation from the sensitized bacteria.

Of all explanations considered, therefore, that of Neisser and Wechsberg seems to be the least likely. It would seem to us that Bordet's interpretation of these facts is borne out indirectly by certain experiments of Morgenroth and Sachs⁵⁷ themselves, in which the mutual quantitative relations between complement and "amboceptor" were studied. In these experiments it was shown that the more highly cells were sensitized, the smaller was the quantity of complement which was needed for their hemolysis, and vice versa, the less the sensitization (the smaller the quantity of amboceptor) the more complement was necessary to produce the same result. The following extract from one of their protocols will illustrate this:

BEEF BLOOD CELLS 5%, 1 C. C., ANTIBEEF GOAT SERUM, GUINEA-PIG COMPLEMENT

| Amount of amboceptor | Relative amount of amboceptor | Amount of complement for complete hemolysis |
|-------------------------|----------------------------------|---|
| .05 | 1 | .008 |
| .2 | 4 | .0025 |
| .4 | 8 | .0014 |

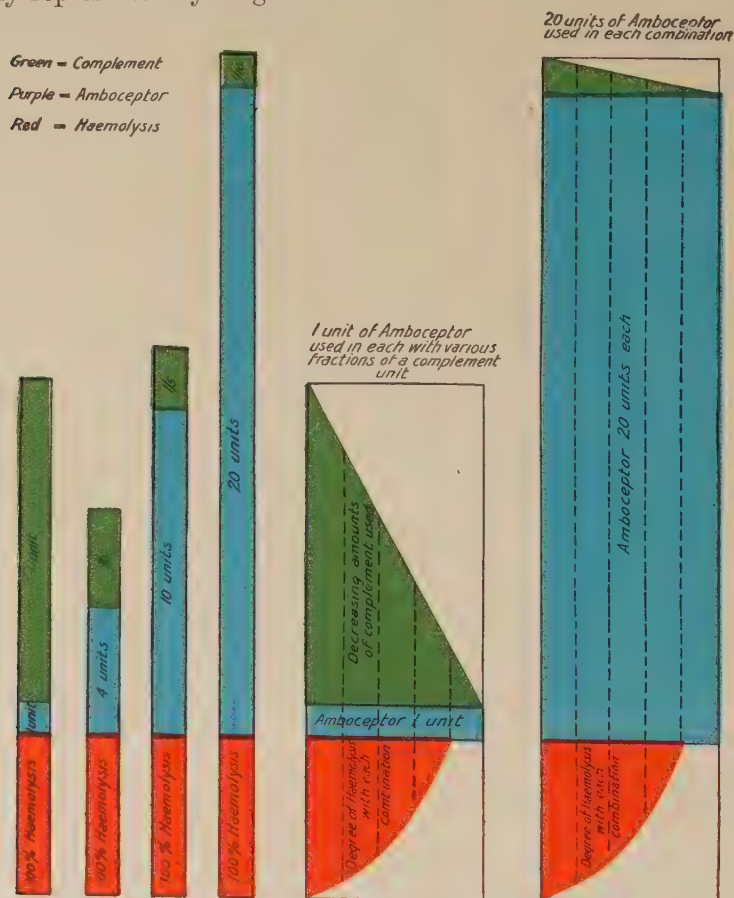
A similar relation may be observed by all who have occasion to work with hemolytic reactions. In the present connection this seems to bear out Bordet's interpretation, since, knowing the differences in functional efficiency of various complements for different hemolytic and bactericidal complexes, we could well expect that insufficient sensitization of a red cell or bacterial antigen, not particularly amenable to the complement employed, might fail to absorb it completely out of the serum, thus giving a negative result which would simulate complete lack of affinity.

This research of Morgenroth and Sachs seems further of funda-

⁵⁷ Morgenroth and Sachs. *Berl. kl. Woch.*, No. 35, 1902.

mental importance in its contradiction of the regularly progressive quantitative relations which strict adherence to the "amboceptor" idea would seem to impose.

The quantitative relations here outlined have been diagrammatically represented by Noguchi as follows:



NOGUCHI'S DIAGRAM ILLUSTRATING THE QUANTITATIVE RELATIONS BETWEEN ANTIGEN, AMBOCEPTOR AND COMPLEMENT.

(Taken from Noguchi, "Serum Diagnosis of Syphilis," Lippincott, Philadelphia, 1910.)

The essential point of difference between the opinions of Ehrlich and Bordet concerning the processes of hemolysis and bacteriolysis lies, as we have seen, in the conception of the union of alexin or complement with amboceptor or sensitizer. Although Ehrlich and his followers admit that the union of complement with amboceptor does not *usually* occur unless the amboceptor has previously united with the antigen, they still maintain that this may occasionally take place

in the case of special complexes in which the complement may directly unite with free amboceptor. This, we have seen, is the basis of the Neisser-Wechsberg conception of complement—"Ablenkung" or deviation, and of other ramifications of this theory. Bordet, on the other hand, consistently holds that alexin or complement is attached only by the complex antigen-sensitizer (antigen-amboceptor). In the controversy which this difference aroused, an observation was reported by Ehrlich and Sachs,⁵⁸ which seemed to represent, as they themselves express it, an "Experimentum Crucis" proving Ehrlich's contention of the intermediary function of the amboceptor in contrast to Bordet's "sensitization" idea. The facts, as they record them, are as follows: When fresh horse serum is added to guinea pig corpuscles, slight hemolysis results. When inactivated ox serum alone is added to such corpuscles, of course no hemolysis results. If the corpuscles are, on the other hand, exposed to the action of the inactive ox serum, together with fresh horse serum, very active hemolysis is brought about. Apparently the ox serum sensitizes (or furnishes amboceptor to) the guinea pig corpuscles, rendering them amenable to the action of the complement in the fresh horse serum. In other words, inactivated ox serum can be reactivated by the addition of fresh horse serum. From this one would expect that if the guinea pig cells were exposed to inactive ox serum, then separated from the serum by centrifugalization and fresh horse serum subsequently added, hemolysis would ensue. However, this was not the case. When the cells were so treated it was found that they had not been sensitized, and, what is more, it could be shown that the ox serum so employed had lost none of its ability to produce strong hemolysis when added to another complex of cells and fresh horse serum. Ehrlich and Sachs concluded that this experiment definitely showed the ability of the amboceptor in the ox serum to unite with alexin independently. The relation to the cell occurred only after the union of the amboceptor in the ox serum and the complement in the horse serum had been established, and if their interpretation is correct, of course, it constitutes strong evidence against the general principle of "sensitization" as conceived by Bordet.

This apparent inability of the corpuscles to absorb amboceptor independently out of the inactivated ox serum, and the fact that hemolysis results only if the corpuscles, ox serum, and fresh horse serum are all simultaneously present, are extraordinary and not at all in keeping with the preceding work of Ehrlich and Morgenroth, and indeed with experience of these phenomena in general. It is logical therefore to examine more closely the peculiar conditions maintained in these experiments before applying the reasoning deduced from obviously different phenomena to their explanation.

⁵⁸ Ehrlich and Sachs. *Berl. klin. Woch.*, No. 21, 1902.

Bordet and Gay⁵⁹ accordingly studied the Ehrlich-Sachs phenomenon carefully and obtained results which confirmed the experimental data of these writers but cast much doubt upon the validity of their conclusions.

In going over the experiments of Ehrlich and Sachs, Bordet and Gay made an observation which had apparently escaped the attention of the former investigators. Heated bovine serum has but a slight agglutinating power for guinea pig corpuscles. Fresh horse serum agglutinates them only slightly and slowly. On the other hand a mixture of the two sera agglutinates them very rapidly and completely. The bovine serum apparently possessed an accelerating or fortifying influence both upon the weakly active normal hemolysins and agglutinins in the horse serum. Bordet and Gay consequently suspected that this property might be due to an undescribed substance, peculiar to the bovine serum. To eliminate the uncertain elements obtaining in experiments in which normal sensitizer is used they now experimented with guinea pig corpuscles, anti-guinea pig sensitizer (from a rabbit immunized with guinea pig blood cells) and guinea pig alexin.

They found that sensitized guinea pig cells are hemolyzed by guinea pig alexin very slowly and imperfectly, as is often the case when the alexin comes from the same animal species as the cells. When heated bovine serum was added to the complex of sensitized cells and alexin, rapid agglutination and hemolysis resulted. Their experiments may be tabulated as follows:

1. Cells + guinea pig alexin + heated bovine serum = no agglutination; very slight hemolysis on next day.
2. Cells + sensitizer + heated bovine serum = slight agglutination; no hemolysis.
3. Cells + sensitizer + alexin + bovine serum = powerful agglutination and complete hemolysis in 10 minutes.
4. Cells + sensitizer + alexin = very slight agglutination and incomplete hemolysis in 30 minutes.
5. Cells + sensitizer = slight agglutination; no hemolysis.

In tube (1) the slight hemolysis was due to the small amount of normal sensitizer present in the bovine serum, and the slight agglutination in tube (5) is referable to the agglutinating power of the sensitizer. In tube (3) we see the powerfully accelerating effects exerted both upon agglutination and hemolysis when bovine serum acts upon sensitized corpuscles in the presence of alexin.

Bordet and Gay's interpretation of the Ehrlich-Sachs phenomenon, in the light of these new experiments then, is, in their own words, as follows: "When guinea pig corpuscles are added to a mixture of the two sera they are affected by the sensitizer of the horse serum and, to a certain extent, by the sensitizer in the heated

⁵⁹ Bordet and Gay. *Ann. de l'Inst. Past.*, Vol. 20, 1906, p. 467.

bovine serum. This second sensitizer is, however, superfluous. Its presence is by no means necessary for the experiment. When this sensitization is effected the corpuscles are then in condition to fix the horse alexin. This alexin, however, has only slight hemolytic power. But once the corpuscles have become sensitized and laden with alexin they are modified in their properties of molecular adhesion to such an extent that they become able to attract a colloidal substance of bovine serum, which unites with them. The adhesion of this new substance produces two results: it causes the blood corpuscles to be more easily destroyed by alexin and also agglutinates them energetically. Consequently, a powerful clumping, followed by hemolysis, is observed."

Bordet and Gay, therefore, assume that the action of the bovine serum is due to a new substance which they speak of as "bovine colloid." This substance resists heating to 56° C., is probably albuminous, and has the property of uniting with cells that are laden with sensitizer and alexin, but remains free in the presence of normal or merely sensitized cells.

They fortify this opinion by showing experimentally that the "colloid" is removed from bovine serum by absorption with sensitized bovine corpuscles which have been treated with horse alexin.⁶⁰

Bordet and Streng⁶¹ later studied this "colloid" more thoroughly and have suggested for it the name "*conglutinin*." Streng⁶² later showed that the agglutinating action of this substance could be shown not only for sensitized and "alexinized" red blood cells, but also for similarly treated bacteria, and that conglutinins were present not only in bovine serum, but in that of goats, sheep, antelopes, and a number of other herbivores, but apparently absent in cats, dogs, guinea pigs, and birds.

The body described by these workers as conglutinin is probably identical with a similar heat-stable serum component reported by Manwaring⁶³ and called by him "*auxilysin*."

⁶⁰ Browning (*Wien kl. Wochenschr.*, 1906) had shown that horse alexin may be absorbed by sensitized beef cells without causing hemolysis.

⁶¹ Bordet and Streng. *Centralbl. f. Bakt.*, Orig. Vol. 49, 1909.

⁶² Streng. *Zeitschr. f. Immunitätsforsch.*, Orig. Vol. 2, 1909, p. 415.

⁶³ Manwaring. *Centralbl. f. Bakt.*, 1906; Orig. Vol. 42.

CHAPTER VII

FURTHER DEVELOPMENT OF OUR KNOWLEDGE CONCERNING COMPLEMENT OR ALEXIN. COMPLEMENT FIXATION

It will be remembered that Buchner in his first studies upon the "alexin" compared its action to that of an enzyme or ferment, and suggested that the source of this substance might possibly be found in the white blood cells. This thought was very obviously suggested by the observation that bacteria were destroyed within the white blood cells, after phagocytosis, by a process analogous in many ways to that by which they were destroyed by the serum constituents. Hankin,¹ in an elaborate study dealing with the problem, maintained the leukocytic origin of alexin on the basis of the observation that increased bactericidal properties closely followed upon the heels of periods of leukocytosis. He assigned the particular property of alexin production to the eosinophile cells, proposing for them the designation "alexocytes." Further study, however, has not justified such an association with the eosinophiles, and Hankin's opinion has not been experimentally upheld.

After Hankin the problem occupied the attention of a number of other investigators, and many of them succeeded in showing that there was, indeed, an increased bactericidal power in exudates rich in leukocytes, and further that bactericidal substances could be directly extracted from leukocytic emulsions. We refer particularly to the early work of Denys and Havet,² of Hahn,³ of Van de Velde,⁴ and others, studies which will be described in our chapter on phagocytosis. This work was done before the complex nature of the bactericidal constituents of serum had been demonstrated and before the work of Schattenfroh and others had shown that the bactericidal substances extracted from leukocytes were of a nature quite distinct from the active elements of the serum, and were independent of the participation of alexin. Although these earlier investigations cannot properly be regarded, therefore, as proving the leukocytic origin of

¹ Hankin. *Centralbl. f. Bakt.*, Vol. 12, 1892.

² Denys and Havet. *La Cellule*, Vol. 10, 1894.

³ Hahn. *Archiv f. Hyg.*, Vol. 25, 1895.

⁴ Van de Velde. *La Cellule*, Vol. 10, 1894.

alexin, Metchnikoff and his school have nevertheless adhered to this conception for various additional reasons.

Metchnikoff distinguishes between two kinds of alexin—the *microcytase*, which is the bactericidal complement or alexin, and is supposed to originate from the microphages or polynuclear leukocytes, and the *macrocytase*, which represents the hemolytic and cytolytic alexin or complement, and originates from the mononuclear cells or macrophages. As in the case of the bactericidal alexin, extraction methods have been employed to demonstrate that the hemolytic alexin took its origin in the macrophages, and at Metchnikoff's suggestion, Tarassewitch⁵ prepared hemolytic substances by extracting spleen tissue and other "macrophagic organs" in various ways. Here again the identity of the hemolytic extracts with serum hemolysins has been placed in doubt. Korschun and Morgenroth⁶ have shown that the hemolytic organ extracts were heat stable and alcohol soluble; Donath and Landsteiner,⁷ and others, have obtained similar results. It would be quite thankless to review the extensive literature which has accumulated upon this point. It would seem, in summarizing it, that no definite proof of the presence of true, active alexin, either hemolytic or bactericidal, within the leukocyte or mononuclear cells has been brought by methods of extraction, and the apparently positive results reported by earlier observers are adequately explained by the discovery of the heat-stable and non-reactivable bactericidal and hemolytic substances in extracts of such cells by Schattenfroh, Korschun and Morgenroth, and many others. It appears, moreover, from these investigations that probably the intracellular substances by which the digestion of ingested bacteria or blood cells is brought about are of a nature entirely distinct from that of the serum antibodies and alexins. A very ingenious demonstration of this is found in an experiment first made by Neufeld. Neufeld⁸ allowed leukocytes to take up highly sensitized red cells. Instead of undergoing prompt hemolysis, as they would if small amounts of alexin had been added, they were slowly broken up without hemolysis, fragments of hemoglobin remaining after complete morphological disintegration of the erythrocytes. At no time were intraphagocytic "shadow" forms observed.

The failure to extract alexins from dead leukocytes does not, however, preclude the possibility of the secretion of alexins by living leukocytes. This point is one which is, of course, much more difficult to investigate directly. Indirectly the increased bactericidal properties of exudates rich in leukocytes, as found by Denys and Havet, would point in this direction. However, even this is not

⁵ Tarassewitch. Cited from Metchnikoff.

⁶ Korschun and Morgenroth. *Berl. kl. Woch.*, No. 37, 1902.

⁷ Donath and Landsteiner. *Wien. kl. Rundschau*, Vol. 40, 1902.

⁸ Neufeld. *Arb. a. d. kais. Gesundheitsamt.*, Vol. 28, 1908, p. 125.

conclusive, since at the time when these investigations were carried out no discrimination was made between the bactericidal serum substances and those other "endolysins" which might well have been extracted from the accumulated white blood cells. The writer some years ago attempted to approach this problem directly by keeping leukocytes alive in inactivated serum and in Ringer's solution at 37.5° C. for several days in the hope that, after 48 hours, alexin, hemolytic or bactericidal, might appear in these fluids. The experiments were entirely negative, but were regarded as inconclusive, since it was impossible to determine accurately how long, or in what proportion, the leukocytes had remained alive.

One of the basic premises of Metchnikoff's theory on the nature of alexin consists in the conception that alexin is not found in the circulating blood plasma, but appears only when there has been leukocytic injury, as in the clotting of blood or in the "phagolysis" which, as we have seen in the chapter on phagocytosis, usually occurs after foreign substances have been injected into the peritoneum, preceding a local accumulation of leukocytes. This point of view seems to be rendered improbable because of the rapid hemolysis which occurs when we inject sensitized red blood cells into the circulation of an animal, but we might here, too, assume a preliminary injury to white blood cells resulting from the intravenous injection.

Much less likely to be accompanied by cell injury is the method of obtaining blood serum by creating an area of artificial edema by ligating a limb—or, as in Metchnikoff's^{9 10} experiments, the ear of a rabbit. And, indeed, in edema fluids so obtained little or no alexin is ordinarily found. This fact has been interpreted in favor of Metchnikoff's views, as has also the curious absence of alexin in the aqueous humor of the anterior chamber of the eye.^{11 12} In this fluid no alexin is present under normal conditions, but if puncture is practiced, and the fluid again taken after a period of three or four hours, alexin is now found, probably, according to Metchnikoff's school, because of the coincident entrance of leukocytes into this space. It is conceivable, however, that the aqueous humor may be free from alexin for other reasons than the absence of leukocytes; and an injury which is followed by the invasion of leukocytes is pretty sure to be followed also by the entrance of the fluid elements of the blood; i. e., alexin.

Much experimental work has been done in which it has been attempted to demonstrate directly that the blood plasma contains no complement or alexin. The most important investigation of this

⁹ Metchnikoff. *Ann. Past.*, Vol. 9, 1895.

¹⁰ Bordet. *Ann. Past.*, Vol. 9, 1895.

¹¹ Metchnikoff. *Loc. cit.*

¹² Mesnil. *Ann. Past.*, Vol. 10.

kind is that carried out by Gengou¹³ in 1901. It was Gengou's primary purpose to obtain the plasma of mammals in such a way that no cell injury would occur. This he accomplished by special methods in which coagulation was avoided without the addition of foreign anticoagulants like hirudin, etc. His technique was, in essence, as follows: He took the blood directly through a paraffined cannula into tubes that had been coated with paraffin, and centrifuged it at low temperatures until cell free. This plasma, taken from the paraffin tubes, quickly clotted, and the material with which the experiments were done actually consisted of blood serum. Upon examining the serum so obtained, he found that it exerted practically no bactericidal action. As a result of this investigation he claims to have demonstrated the truth of Metchnikoff's contention that the circulating blood plasma contains no alexin.

If borne out, it is true that Gengou's results would very powerfully support this theory, and for this reason a large number of experiments have been made since then, with the same end in view.

In all such investigations the technical procedures are extremely difficult and, as Addis¹⁴ has recently said, in our opinion quite correctly, it would be impossible to carry out bacteriolytic or hemolytic experiments with mammalian paraffin plasma without obtaining coagulation, and for this reason most of the writers who have repeated Gengou's experiments have worked, as did he, not with plasma, but with serum. Falloise,¹⁵ following Gengou's method exactly, obtained results diametrically opposed to those of Gengou; Schneider,¹⁶ also with the same technique, failed to confirm Gengou's results; Herman,¹⁷ on the other hand, confirms Gengou.

In order to overcome the technical difficulties encountered in working with mammalian plasma a number of writers have more recently experimented with bird blood, which, as is well known, coagulates much more easily than does mammalian blood. Hewlett,¹⁸ who worked with goose plasma and peptone plasma, could not confirm Gengou's results. Lambotte,¹⁹ examining the plasma of chickens, found no difference between the serum and plasma in their contents of bactericidal alexin, as measured against cholera spirilla. Von Dungern, working with fish plasma, obtained similarly negative results, and recently Addis, in a careful comparative study of chicken plasma, found no evidence of differences between plasma and serum in either the bactericidal or the hemolytic alexin. As far

¹³ Gengou. *Ann. de l'Inst. Past.*, Vol. 15, 1901.

¹⁴ Addis. *Journ. of Inf. Dis.*, Vol. 10, 1912.

¹⁵ Falloise. *Bull. de l'Acad. Roy. de Méd.*, 1905, p. 230.

¹⁶ Schneider. *Archiv f. Hyg.*, 1908, Vol. 65, p. 305.

¹⁷ Herman. *Bull. de l'Acad. Roy. de Méd.*, 1904, p. 157.

¹⁸ Hewlett. *Archiv f. exp. Path. u. Pharmk.*, 1903, Vol. 49, p. 307.

¹⁹ Lambotte. *Centralbl. f. Bakt.*, I, Orig., 1903, Vol. 34, p. 453.

as we can tell at present, therefore, we cannot accept, as conclusively proven, the contention that the circulating plasma contains no alexin. Nevertheless the Metchnikoff school have not been discouraged by the various contradictions of Gengou's work, found in the experiments we have enumerated, because they are not satisfied that the technique of other workers has conclusively excluded cell injury. Owing to the great difficulties of investigations of this kind, when carried out with mammalian blood, it is not impossible that they are justified in this, but nevertheless the assumption of the absence of alexin in the plasma finds so many objections in other observations that the burden of proof would certainly rest with Gengou and his supporters. Not the least important of these objections, it seems to us, is based on the very simple experiment of injecting bacteria into the veins of a living animal and finding a very rapid and active phagocytosis. And considering the very probable participation of alexin in the opsonic functions this would seem to point strongly toward the presence of these substances in the circulating blood. The evidence also furnished by the recent developments of our understanding of anaphylaxis would further tend to strengthen our belief in the presence of alexin or complement in the normal circulation. For, in the process, as we shall see in a later chapter, complement plays an important rôle. When 3 per cent. salt solution is administered (as in Friedberger's experiments), and the action of complement is thereby inhibited, anaphylactic shock may be greatly diminished.

It has also been claimed, chiefly by Walker²⁰ and by Henderson Smith,²¹ that, as serum stands upon the clot it at first gains in alexin or complement contents, an occurrence which they attribute to the liberation of alexin from the leukocytes. This observation has not been universally borne out and, even were it unquestionable, it might be dependent upon any one of the numerous factors involved in the complicated process of coagulation rather than upon leukocytic changes only.

The failure to obtain definite proof of the origin of alexin from the white blood cells has led to search for the source of these substances in various organs. An interesting series of investigations on this subject are those of Mlle. Louise Fassin,²² who believes that she has found reasons for definitely associating the thyroid gland with alexin production. She found that the subcutaneous injection of thyroid extract into dogs and rabbits was followed by a rapid increase of alexin, both hemolytic and bactericidal, and that the same thing was true when thyroid substance was administered by mouth. When the thyroid gland was removed from rabbits a reduction of alexin resulted. Although important, these researches do not

²⁰ Walker. *Journ. of Hyg.*, Vol. 3, 1903.

²¹ Smith. *Proc. Roy. Soc.*, Series B, Vol. 79, 1906.

²² Louise Fassin. *C. R. de Soc. Biol.*, Vol. 62, 1907.

necessarily prove that the thyroid can be looked upon as a source of alexin, and, indeed, Fassin gives experimental results without drawing any very sweeping conclusions. It might well be that the thyroid secretion is simply concerned in stimulating the production of alexin from another source. Marbé²³ has similarly associated the thyroid gland with the production of opsonins, which, when we consider the probable identity of alexin and normal opsonin, may be taken as a confirmation of Fassin's work.

Of great interest also are the series of investigations which associate the liver with the production of alexin. The basis of such investigations is found in the observations made by Morgenroth and Ehrlich²⁴ that there is a diminished production of complement or alexin in dogs subjected to phosphorus poisoning, with consequent degeneration of the liver. The first investigator to study this question experimentally was Nolf.²⁵ Nolf tried to approach it by extirpating the liver in dogs, and found that his results were unreliable by this method. He then experimented with rabbits and found that when the liver was extirpated in these animals and the vena cava anastomosed with the portal vein (Eck fistula) the animals would survive for three or four hours. This period, though short, was sufficient to show definite changes in the blood. Taken just before death it differed from that taken just before the operation in a number of important respects. There was relative incoagulability, there was autohemolysis, and with these there occurred an extreme fall of alexin or complement. Serious objections may be brought against Nolf's experiments. In the first place the operation as performed by him results in shock and injury so profound that rapid death ensues, conditions under which not only the complement-producing functions but all functions, secretory and otherwise, are reduced. Müller²⁶ objects to Nolf's experiments chiefly for the reason that he did not prevent the absorption of toxic substances from the intestine, materials which could now enter the general circulation without any longer being neutralized by the liver functions. Müller, for this reason, repeated Nolf's work but, by a complicated technique, temporarily shut off the intestinal circulation in addition to extirpation of the liver. He found, in agreement with Nolf, that exclusion of the liver from the circulation resulted in the prompt diminution of complement or alexin. In all such experiments, however, the very profound shock which necessarily occurs in the animals would seem to us to vitiate the results. Moreover, Liefmann²⁷ has repeated Müller's experiments without being able to obtain the same results. Not satisfied

²³ Marbé. *C. R. de la Soc. Biol.*, Vols. 64, et seq., 1908-1909.

²⁴ Morgenroth and Ehrlich. In Ehrlich's "Gesammelte Arb.," etc.

²⁵ Nolf. *Bull. de l'Acad. de Science de Belg.*, 1908.

²⁶ Müller. *Centralbl. f. Bakt.*, Vol. 57, 1911.

²⁷ Liefmann. *Weichhart's Jahresbericht*, Vol. 8, 1912, p. 155.

with these experiments, however, Liefmann experimented on frogs; in whom, as Friedberger has shown, extirpation of the liver is not so rapidly fatal as in warm-blooded animals. He removed the livers of frogs in a number of cases and, although his animals lived about a week, there was no definite diminution of the hemolytic properties of the serum. It seems, therefore, that the origin of alexin in the body is by no means settled and requires further investigation.

Equally unsatisfactory have been the attempts to define the chemical nature of the complement or alexin. In the investigations dealing with the hemolytic action of cobra venom it seemed at first as though a clue to this problem had been found. Flexner and Noguchi²⁸ made the interesting observation that cobra poison alone does not hemolyze the blood cells of certain animals, namely those of cattle, goats, or sheep, if these cells are washed entirely free of serum. This seemed to suggest that the serum of these animals contained some activating substance. It also seemed to indicate that the cells of other animals, which were easily hemolyzed, even when entirely freed of serum, might contain such an activating substance within themselves. The behavior of this activating substance toward snake-venom hemolysis was therefore very similar to the action of complement, except in one important respect, namely, as Calmette²⁹ showed, almost all sera were rendered more efficient for the activation of snake venom when heated to 65° C., whereas complementary properties of sera for other hemolyzing complexes are, of course, destroyed at 56° C. Kyes,^{30 31} on further studying these phenomena, extracted the red blood cells of rabbits and other animals whose cells were hemolyzed by snake venom alone, by shaking them up with distilled water, and showed that, with these extracts, he could activate the venom against ox, goat, and sheep corpuscles, cells which were not ordinarily hemolyzed by the venom without the addition of serum. Similar activation of the venom with extracts of the ox, goat, or sheep corpuscles was not possible. He concluded from this that the blood cells of the rabbit, dog, guinea pig, and man possessed an "endocomplement" for the snake venom; that is, a complementary substance contained within the cells, while in the other species it was found in the activating serum only.

The thermostability of such venom "complements" encouraged him to attempt their isolation, and he found that they were ether-soluble, indicating their lipoidal nature; and, finally, after several negative attempts with activation by other lipoids, he determined that lecithin, added to the corpuscles and the snake venom, brought

²⁸ Flexner and Noguchi. *Journ. Exp. Med.*, Vol. 6, 1902; *Univ. Pa. Med. Bull.*, 1902 and 1903.

²⁹ Calmette. *C. R. de l'Acad. des Sciences*, p. 134, 1902.

³⁰ Kyes. *Berl. klin. Woch.*, Nos. 38 and 39, 1902.

³¹ Kyes and Sachs. *Berl. klin. Woch.*, Nos. 2-4, 1903.

about a rapid hemolysis. This seemed to explain both why heated serum could activate the venom in some cases, and why some varieties of blood cells could be hemolyzed without serum, since lecithin is a substance widely distributed both in the fluids and cells of the animal body. His further studies seemed to show that, by proper chemical manipulation (bringing together cobra poison with lecithin in chloroform solution), he could produce a combination of the two which he called "cobra lecithid," a substance which apparently "activated" cobra venom. He conceived it as the "amboceptor-complement" complex of the cobra hemolysin, which acted hemolytically upon all varieties of blood cells.

These researches of Kyes aroused much interest, chiefly because they seemed to furnish an example of a chemically definable complement, lipoidal in its constitution. Recent researches by Von Dungern and Coca,³² however, seem to prove that, while Kyes' experimental facts were perfectly accurate, his conclusions do not seem to have been warranted. Von Dungern and Coca showed that the cobra venom contains a lipid-splitting ferment which acts upon the lecithin, liberating substances from it which hemolyze in the same way as do many other non-specific substances. The cobra-lecithid, according to this, would represent merely a lecithin derivative which happens to have hemolytic action without any specific relationship to the hemolytic properties of the venom itself. Thus, even in this case, unfortunately, we are not in possession of facts which bring us nearer to a chemical understanding of the complementary substances or alexins.

In the further development of attempts to define alexin or complement chemically, two further researches are of importance, namely, those of Von Liebermann³³ and of Noguchi. In both investigations it is suggested that the alexin may consist of a combination of soaps and proteins. Noguchi³⁴ showed that the hemolytic organ-extracts described by various observers were soaps, a possibility which had been previously considered by Sachs and Kyes.³⁵ Noguchi further established analogies between his soaps and complement as follows: Sensitized blood cells are hemolyzed by mixtures of soaps and inactivated guinea pig serum, while normal erythrocytes are not hemolyzed by similar mixtures; furthermore, like normal complement, such serum-soap mixtures are inactivated by prolonged preservation and by heating at 56° C. Objections were soon made to the findings of both Noguchi and Von Liebermann by Hecker,³⁶ whose experiments seemed to show that when sensitized

³² Von Dungern and Coca. *Münch. med. Woch.*, 1907, p. 2317.

³³ Von Liebermann. *Biochem. Zeitschr.*, Vol. 4, 1907.

³⁴ Noguchi. *Biochem. Zeitschr.*, Vol. 6, 1907.

³⁵ Sachs and Kyes. *Berl. kl. Woch.*, 2-4, 1903.

³⁶ Hecker. *Arb. auf dem könig. Inst. f. exp. Ther.*, Heft 3, 1907.

blood cells were thoroughly washed free of serum soaps did not have this hemolyzing action, and Friedemann and Sachs³⁷ claimed that they were unable in any case to inactivate the hemolytic serum soap mixtures by heating to 56° C. These writers, as well as others, attribute Noguchi's results to the fact that the sera which he used to produce his "artificial complement," i. e., his serum soap mixtures, were heated to 50-51° C. only, a fact which would justify doubt of complete inactivation. Knaff-Lenz³⁸ has more recently carried out experiments on the same question. His results seem to show that the hemolytic action exerted by fatty acids or soaps is a phenomenon quite incomparable to true complement action, and that these hemolysins are heat stable, remaining unchanged by heating at 56° C. We have referred in a number of places to the analogy between alexins and ferments or enzymes. The chief objection to this conception formerly brought forward was based upon the fact that the complement or alexin, unlike an enzyme, was used up during its reactions, and that a definite quantitative relationship existed between the alexin and the amount of cells or bacteria upon which it could act. Recent experiments by Kiss³⁹ seem to show that this quantitative relationship is not as strict and regular as was formerly supposed. He showed that the action of complement depends very largely upon its concentration. For instance, to cite his work directly: "0.05 complement is sufficient to hemolyze completely a definite quantity of sensitized blood cells if the experiment is done in a total volume of 5 c. c. 0.02 c. c. of complement gives absolutely no hemolysis in a similar volume. When, however, the total volume is reduced to 2.5 c. c., then 0.02 c. c. of the complement begins to act, and it produces complete hemolysis if the total volume is reduced to 1.25." In further developing this observation he showed that, if sufficiently concentrated, a very small amount of complement can act upon an extremely large amount of red blood cells, an amount incomparably larger than those acted upon in more dilute solutions. These observations would tend to strengthen considerably the conception of the ferment nature of alexins in general.

Kiss' observations are furthermore in agreement with the investigations of Liefmann and Cohn,⁴⁰ whose work we have mentioned in the preceding chapter on Cytolysis. These writers assert that the fixation of complement during hemolysis is not due to its chemical union with the sensitized cells, but is due to fixation by the end products of the reaction; in other words, by the stromata of the red cells and possibly by other substances given up by these cells. A further factor contributing to the disappearance of complement in

³⁷ Friedemann and Sachs. *Biochem. Zeitschr.*, Vol. 12, 1908.

³⁸ Knaff-Lenz. *Biochem. Zeitschr.*, Vol. 20, 1909.

³⁹ Kiss. *Zeitschr. f. Imm.*, Vol. 3, 1909.

⁴⁰ Liefmann and Cohn. *Zeitschr. f. Imm.*, Vol. 8, 1911.

such reactions is, they claim, its rapid deterioration at 37° to 40° C., when diluted. If they are right, these considerations also remove important objections to the conception of complement as a ferment.

It is clear, therefore, that although we have gained much detailed information regarding the functional activity of the complement or alexin, and may assume, in a general way, that its action is similar to, if not identical with, that of an enzyme, we are nevertheless still very much in the dark concerning its chemical nature. The same thing may be said in regard to its physical characteristics. One method of investigating the physical properties of complement has been that of filtration. It may be remembered that one of Ehrlich and Morgenroth's⁴¹ arguments in favor of the multiplicity of complement was the fact that, when goat serum was filtered through a Pukal candle, the complement which was active upon rabbit corpuscles was retained, while that which acted upon guinea-pig cells passed through. Immune bodies or amboceptor always passed through.

Vedder,⁴² in similar experiments upon bactericidal complements, claims to have been able, in the same way, to separate the complements acting upon different bacteria. The problem has been more recently investigated by Muir and Browning.⁴³ Their conclusions are briefly as follows: In the early stages of filtration through a Berkefeldt filter complement is often completely held back. After continued filtration it begins to pass through. If the complement is inactivated by the addition of hypertonic salt solution (5 per cent.), it passes through, and the filtrate can be reactivated by dilution to isotonicity. Sensitizer or amboceptor always passes through. Just how these experiments are to be interpreted is a little obscure. The fact that the addition of salt renders the complement capable of passing through the filter would seem to indicate that its original inability to permeate did not depend upon the size of the molecule. On the other hand, it is also possible that the addition of salt to the complement may increase its dispersion in such a way that the individual particles are rendered smaller. This, however, is purely speculative, and we are at a loss for a fully satisfactory explanation of the results of Muir and Browning. We have repeated some of the experiments of Muir and Browning and, in substance, confirmed their results. It is our opinion that new filters remove complement by adsorption, just as this is accomplished when complement is shaken up with kaolin or other finely suspended material.

That the addition of salts of various kinds in quantities greater than isotonicity (or more than the equivalent of $0.85=0.9$ per cent.

⁴¹ Morgenroth and Ehrlich. Ehrlich's "Gesammelte Arbeiten," etc.

⁴² Vedder. *Journ. Med. Res.*, Vol. 9, 1903.

⁴³ Muir and Browning. *Journ. of Path. and Bact.*, Vol. 13, 1909.

NaCl)⁴⁴ exerts a profound action upon the activity of complement is well known. Nolf⁴⁵ noted this in 1900, and the problem has been studied since that time by many investigators. Von Lingelsheim,⁴⁶ who studied it in connection with his work on the refutation of the "osmotic" theories of immunity, showed that increasing the salt contents of serum (KNO_3 , NaCl, K_2HPO_3 , etc.) progressively diminished its bactericidal power. Hektoen and Ruediger⁴⁷ also, after a very thorough study of this phenomenon, conclude that the action of the salts in such cases is exerted upon the alexin or complement and not upon the heat-stable sensitizers, and that it probably depends upon "physicochemical" causes. However, the manner in which such salt-inactivation is brought about is, to a great extent, obscure. There is no visible precipitation from serum after the addition of salts sufficient in quantity to weaken its action. Nothing is, as far as we can tell, removed from solution, and yet there is temporary inactivation which, at the same time, renders the complement filterable, facts from which we can only surmise some physical alteration.

Inactivation of the complement also follows the removal of salts, but here the process is accompanied by a definite chemical change in that the serum globulins are precipitated.

Studies of this process have led to important modifications in our conception of the nature of alexin, since they have shown that this body, formerly assumed to be single and homogeneous, may be subdivided into at least two component parts by a number of experimental procedures. Ferrata was the first one to point this out as a consequence of investigations undertaken by him primarily with the purpose of determining the nature of the influence of salts upon hemolytic processes. Older studies of Buchner and Orthenberger⁴⁸ had shown that bactericidal action was inhibited when salts were removed from the medium, but the causes underlying such inhibition had not been made clear. Ferrata⁴⁹ found, in the first place, that the absence of salts exerted no effect upon the mechanism of sensitization, but that amboceptor or sensitizer became attached to the cellular elements as readily when salts were absent as when the reagents were suspended in normal salt solution. It was a natural inference, therefore, that the failure of hemolysis, which he observed in salt-free media (analogous to the similar experiences of Buchner

⁴⁴ Alexin can be preserved in the refrigerator for long periods if hypertonic salt solution (15 to 25%) is added. It will again become active if isotonicity is restored with distilled water.

⁴⁵ Nolf. *Ann. Past.*, Vol. 14, 1900.

⁴⁶ V. Lingelsheim. *Zeitschr. f. Hyg.*, Vol. 37, 1901.

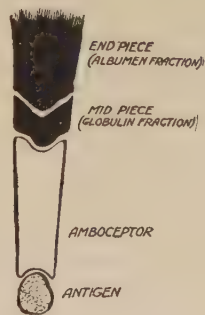
⁴⁷ Hektoen and Ruediger. *Journ. of Inf. Dis.*, Vol. 1, 1904.

⁴⁸ Buchner and Orthenberger. *Archiv f. Hyg.*, Vol. 10, 1890.

⁴⁹ Ferrata. *Berl. kl. Woch.*, 1907, No. 13.

in the case of bacteriolysis), must be attributed to failure of functionation on the part of the complement. On further investigation he obtained a very simple explanation. Ferrata removed the salts from his sera by dialyzing for twenty-four hours against distilled water. In this process, of course, there is a precipitation of the globulins while the water-soluble albumins remain in solution. The former may be redissolved in normal salt solution and the latter rendered isotonic by the addition of calculated amounts of concentrated salt. In this way the original serum components are divided into two parts, neither of which, as Ferrata found, is alone capable of producing hemolysis of sensitized cells. In order to obtain the complementary action possessed by the original serum it is necessary to combine the two. This principle discovered by Ferrata is probably responsible also for the results obtained by Sachs and Terunchi,⁵⁰ who likewise noted the destruction of the complementary function in sera diluted with distilled water, but attributed this, in their publication, to the action of a complement-destroying ferment, which is assumed to be active in salt-free media only.

In his first experiments Ferrata reported that the precipitated globulin fraction was thermostable, the thermolability of complement being due entirely to the unprecipitated albumin fraction. The work of Ferrata was soon continued, however, by a number of other workers, who confirmed the essential fact of the partition of the complement but modified and considerably extended the original observations. Brand⁵¹ found that both fractions were equally thermolabile, and that the globulin sediment, after being redissolved in salt solution, could not be preserved in an active condition for more than a few hours. Preserved in distilled water or as sediment, it may retain its activity for several days, but dissolved in salt solution it becomes inactive within 3 to 4 hours, at room temperature. Michaelis and Skwinsky⁵² have since shown that the globulin fraction, thermolabile when free, is unaffected by a temperature of 56° C. after it has become attached to sensitized cells. Brand further studied the relationship of the two fractions to the sensitized cells and found that the globulin fraction may attach directly to such antigen-antibody complexes, but that the albumin fraction cannot be bound in this way unless the globulin fraction has been previously attached. For this reason he has referred to the former as the "end-



CONCEPTION OF COMPLEMENT-SPLITTING AS FIRST SUGGESTED BY BRAND.

⁵⁰ Sachs and Terunchi. *Berl. kl. Woch.*, 1907, Nos. 16, 17, and 19.

⁵¹ Brand. *Berl. kl. Woch.*, 1907, No. 34.

⁵² Michaelis and Skwinsky. *Zeitschr. f. Imm.*, Vol. 4, 1910.

piece" and the latter globulin sediment as the "mid-piece," assuming, on the basis of the conception of Ehrlich, that the globulin fraction serves to establish a link between the sensitized cell and the end-piece analogous to that formed by the "amboceptor" between the cell and the whole complement. It is possible, therefore, to treat sensitized cells with mid-piece in such a way that they are thereafter susceptible to hemolysis by the end-piece alone. Such cell-sensitizer-mid-piece combinations have been spoken of by Michaelis as "*persensitized*" cells.

Tsurusaki⁵³ confirmed the findings of Brand as to the thermostability of both "mid-piece" and "end-piece," but was unable to separate the complement of *normal* hemolysins into the two components in the same way, since he found that hemolytic power was, in such cases, completely destroyed after twenty-four hours of dialysis. It seems to us not impossible that the natural deterioration of alexic power which takes place during such periods of time, at temperatures of from 16° to 20° C., may easily be held accountable for this, since the very feeble sensitization of cells in normal hemolysin complexes requires a correspondingly larger amount of alexin for activation.

We have mentioned that the so-called "mid-piece" undergoes a rapid change when dissolved in salt solution and, after 3 or 4 hours, may lose its ability to induce hemolysis when added to sensitized cells together with end-piece. Although Hecker⁵⁴ was able to confirm this, he nevertheless showed that this fact does not imply a destruction of the mid-piece. For when such apparently inactive "mid-piece" was added separately to sensitized cells, and end-piece was subsequently allowed to act upon the complex, hemolysis resulted. This seems to show that the "mid-piece" undergoes a change on standing in salt solution which does not alter its ability to combine with the sensitized cells, but which subjects it to inhibition of such union when end-piece is present. It is also a peculiar fact, evident in many of our own experiments, that when "mid-piece" and "end-piece" are first mixed and then added to sensitized cells the effect in hemolysis is less powerful than when the "mid-piece" is added to the cells first, and the "end-piece" later. This effect is so instantaneous that, if, in a series of experiments in which combinations of mid- and end-piece are used, the end-piece is run into the tubes containing the cells just before the mid-piece is added instead of the other way round, hemolysis is inhibited.

In working with dialysis, also, we have regularly had an experience which may explain the difficulties which many other investigators have had in such experiments. The globulin precipitate, which

⁵³ Tsurusaki. *Biochem. Zeitschr.*, Vol. 10, 1908.

⁵⁴ Hecker. *Arb. a. d. könig. Inst. f. exp. Ther.*, Frankfurt a/M., Heft 3, 1907. See also Guggenheimer. *Zeitschr. f. Imm.*, Vol. 8, 1911.

fell out after dialysis of 24 hours or more, almost without exception, retained moderate or slight hemolytic properties, which could not be removed until the precipitate had been dissolved in salt solution and reprecipitated with distilled water two or three times. This would imply that a minute amount of the end-piece, carried down in precipitation, must suffice to activate the mid-piece and would seem to point to the fact that in whole serum the two fractions are present as a complex and not separately. This question has been much discussed and many facts have been brought out on both sides. Hecker showed that the combination of mid-piece with the sensitized cells can take place at a temperature of 0° C., while that of end-piece with the "persensitized" cells requires a considerably higher temperature. The bearing this fact may have upon similar earlier experiments of Ehrlich and Morgenroth upon the thermal conditions governing the union of amboceptor and complement with antigen is self-evident. In the present connection, however, the fact that the two fractions may be separately absorbed out of the serum by sensitized cells at 0° C. would suggest the probability of their being separate in the whole blood. No crucial experiment has so far been possible, and there is not enough evidence on either side as yet to justify a definite opinion. However, the experiments of Michaelis and Skwirsky and later ones of Skwirsky alone have much indirect bearing on this question, though final interpretation is as yet impossible. Michaelis and Skwirsky,⁵⁵ after determining that an acid reaction inhibits the hemolysis of sensitized blood cells, found that under such conditions "mid-piece" alone is bound, but that "end-piece" or the albumin fraction is left unbound. They recommend the use of strongly sensitized cells in an acid medium as a method of obtaining free "end-piece" from serum.

Skwirsky⁵⁶ subsequently found that during the ordinary Wassermann reaction the complex of syphilitic serum and antigen binds the mid-piece only. If the Wassermann reaction has been strongly positive; that is if there has been absolutely no hemolysis, and we remove the supernatant fluid by centrifugation, active end-piece can be demonstrated in it by the addition of persensitized cells. Bronfenbrenner and Noguchi have also studied this phenomenon, but do not believe that Skwirsky's experiments prove that end-piece is free in such "fixation" supernatant fluids. These supernatant fluids, according to them, differ from all other "end-pieces" in that they are active upon persensitized sheep corpuscles only, but not upon other cells. An explanation for this is lacking.

There is much that is confusing in the facts so far revealed about the two component parts of the alexin. The most difficult fact to explain is the peculiar inactivation of the mid-piece in salt solution,

⁵⁵ Michaelis and Skwirsky. *Zeitschr. f. Imm.*, Vol. 4, 1910.

⁵⁶ Skwirsky. *Zeitschr. f. Imm.*, Vol. 5, 1910.

which prevents its functionation in the simultaneous presence of end-piece, but does not seem to interfere with its ability to combine with the sensitized cells. As was to be expected, explanation for this has been sought by the Ehrlich school in changes of affinity. Sachs suggests that the mid-piece, by its preservation in salt solution, has lost its avidity for the sensitized cells and has gained in avidity for the end-piece, an alteration which therefore prevents its union with the cells. The same idea was suggested by Hecker himself. It is a little difficult to reconcile this explanation, however, with the fact that whole serum can be preserved and remain active in its complementary function for a number of days, mid-piece and end-piece being present together, in a medium which, as far as salt contents are concerned, is isotonic with the salt solution in which mid-piece deteriorates so rapidly when alone.

That there is, after all, much similarity between the alexins of different animals is evident from the fact that, as Marks and others have shown, the end-piece of one animal may activate the mid-piece of another species. It appears also from experiments like those of Ritz and Sachs⁵⁷ that an animal may possess a mid-piece for certain sensitized cell complexes without possessing a corresponding end-piece. Thus they found that the serum of mice contained a mid-piece but not an end-piece for sensitized guinea pig corpuscles.

Much that has been found out about the so-called globulin portion, moreover, tends to engender doubt as to the wisdom of applying to these complement fractions the terms "mid-piece" and "end-piece," an objection which is based upon reasons similar to those which prevent Bordet from accepting the term amboceptor. For so little is actually known concerning the mechanism of complement functionation, that it seems unwise to establish on a firm basis a preconceived idea of the mechanism by adapting the terminology to a theory. The most confusing feature of the problem lies in the surprising quantitative relations which seem to exist in the reactions of the two fractions. Thus Liefmann and Cohn⁵⁸ claim that in the presence of moderately sensitized cells no measurable amount of the so-called mid-piece or globulin fraction is bound, that is, removed from solution; and yet, when both fractions are added to such cells, rapid and complete hemolysis results. In the presence of heavily sensitized cells (20 to 50 units) a small quantity only is removed. Nevertheless this fraction has had a demonstrable effect on the cells, since it has rendered them amenable to the action of the albumin fraction. In all such experiments, therefore, as Liefmann justly points out, the degree of sensitization must be taken into consideration before conclusions are formulated. It is curious also that a slight excess of the globulin fraction may prevent complement action completely. In

⁵⁷ Ritz and Sachs. *Zeitschr. f. Imm.*, Vol. 14, 1912.

⁵⁸ Liefmann and Cohn. *Zeitschr. f. Imm.*, Vol. 7, 1910.

experiments cited by Marks⁵⁹ it appears that the most ineffective complement is obtained when "mid-piece" and "end-piece" are added to the sensitized cells in proportions of 1 to 1. If the proportion of "mid-piece" is increased two or threefold over that of "end-piece," hemolysis is inhibited. This, however, is true only when the two fractions are simultaneously added to the sensitized cells. When the sensitized cells are exposed to the excessive quantity of the "mid-piece" separately, and "end-piece" added later, the effect is one of stronger hemolysis than when smaller amounts are used. It is thus seen that the relations between the complement fractions in hemolysis are very involved. All that we can be sure of is that there are at least two separable parts, that one of these acts directly upon the sensitized cells, forming a so-called persensitized complex and rendering them amenable to the subsequent action of the unprecipitated albumin fraction.

The many difficulties encountered in the interpretation of the confusing phenomena observed in connection with this problem have, very naturally, led to a corresponding multiplicity of opinion. Most observers at present incline to the opinion that the globulin and albumin portions of fresh serum, separated by Ferrata's or any other of several common methods, represent actually two complement fractions. This is not, however, accepted by all workers. Bronfenbrenner and Noguchi⁶⁰ believe that the entire active complement is contained in the albumin fraction or so-called "end-piece." They hold that "complement-splitting" by dialysis or other methods is an inactivation of end-piece by change of reaction. In their experiments they were able to restore the functional activity of end-piece by the adjustment of reaction, either with acid or alkali, respectively, or by the addition of amphoteric substances. The mid-piece activates, they believe, by reason of its amphoteric nature and consequently adjusts any excessive acidity or alkalinity of the medium. They were able to substitute for mid-piece indifferent amphoteric substances such as alanin. Liefmann⁶¹ has been unable to confirm the experiments of Bronfenbrenner and Noguchi, and believes that their results were caused by incomplete splitting of the complement. Incidental to a study of normal opsonins the writer has also repeated the experiments of Bronfenbrenner without being able to confirm them.⁶²

The method of Ferrata for the separation of the two parts of the complement is successful only if dialysis is very thorough and sufficiently prolonged to lead to complete precipitation of the globulins. Neufeld and Haendel⁶³ have had difficulty in thus separating the

⁵⁹ Marks. *Zeitschr. f. Imm.*, Vols. 8 and 11, 1911.

⁶⁰ Bronfenbrenner and Noguchi. *Jour. of Exp. Med.*, Vol. 15, 1912.

⁶¹ Liefmann. *Weichhardt's Jahresbericht*, Vol. 8, 1912.

⁶² Zinsser and Cary. *Journ. of Exp. Med.*, Vol. 19, 1914.

⁶³ Neufeld and Haendel. *Arb. a. d. kais. Gesund.*, 1908.

fractions, and the writer has noticed similar failures but has always been able to obtain eventual separation by sufficient prolongation of the dialysis. Because of the occasional difficulties and because of the time-consuming and inconvenient nature of the method other means of separation have been devised. The one used with success by many workers has been that introduced by Sachs and Altmann,⁶⁴ namely, precipitation of the sera with weak hydrochloric acid $\frac{N}{360}$ to $\frac{N}{250}$. Liefmann has separated the components by precipitation of the globulins by the introduction of CO_2 . In carrying out this method, Fraenkel⁶⁵ has found it advantageous to dilute the serum ten times with distilled water, then allowing the CO_2 to flow in at low temperatures. It is likely that any of the usual methods of globulin separation will serve for complement partition. The salting out methods are, however, extremely inconvenient because of the prolonged dialysis subsequently necessary to remove the salts.

The inactivation of complement or alexin by the addition of salts or by splitting is, very apparently, a temporary inactivation in which prompt restitution can be practiced by bringing back original conditions either by dilution to isotonicity or by reconstruction of the divided substance, respectively. Heating to 56°C ., the simplest and most commonly employed method of inactivations was, until of late, regarded as an irreversible process, the complement being irretrievably destroyed in the procedure. Gramenitski⁶⁶ has recently carried out experiments which seem to show that this opinion is erroneous. His experiments were suggested by the fact, observed by Bach and Chodat,⁶⁷ that certain oxydases and diastases may spontaneously regain some of their activity after inactivation by heat. His work with complement indicated a similar gradual return to an active condition after moderate heating. The great theoretical importance of this observation will justify our insertion of one of Gramenitski's protocols.

Experiment 1. Complement 10 times diluted was heated to 56°C . for 7 minutes. It was then tested against sensitized beef blood at varying intervals as follows:

| Time after heating at which test was made | Quantity of hemoglobin gone into solution after | | | |
|---|---|-----------|-----------|-----------|
| | % 10 min. | % 20 min. | % 30 min. | % 40 min. |
| Immediately after heating..... | 0 | 20 | 40 | 70 |
| 1½ hour..... | 0 | 30 | 60 | 80 |
| 24 hours..... | 20 | 70 | 80 | 100 |
| 48 hours..... | 10 | 40 | 70 | |

⁶⁴ Sachs and Altmann. Cited from Sachs in "Kolle u. Wassermann Handbuch," Vol. 2, p. 877.

⁶⁵ Fraenkel. *Zeitschr. f. Imm.*, I, Vol. 8, 1911.

⁶⁶ Gramenitski. *Biochem. Zeits.*, Vol. 38, 1912.

⁶⁷ Bach and Chodat. Cited from Gramenitski, *loc. cit.*, p. 511.

In other experiments in which heating was more prolonged a similar regeneration was observed, though not as pronounced as in the one cited above. The largest amount of restored complement seemed to be present after about 24 hours. After this gradual deterioration again ensued. It is quite impossible to offer an adequate explanation for this at the present time. Gramenitski⁶⁸ acknowledges this, but permits himself certain speculations which we repeat in nearly his own language, since there is much in them which seems to us reasonable. The complement, as indeed all other active serum constituents, must be looked upon as colloidal in nature. When heat is applied to such substances alterations occur which gradually lead to coagulation. As this occurs there is an aggregation of particles and a consequent diminution of surface tension. This last point has been experimentally demonstrated by Traube,⁶⁹ who has regularly found a fall of surface tension as serum was heated to 56° C. And of greatest interest in this connection is the further determination by Traube that a gradual restoration of the surface tension takes place as the serum is allowed to stand. It is not inconceivable, therefore, that the inactivation of complement by heat may depend upon an alteration of its colloidal state, i. e., an aggregation of the particles, which, if not carried too far, may be reversible and followed by a gradual dispersion as the serum is kept 24 hours. On the same grounds the gradual deterioration of complement on standing may be compared to the slow settling out of colloidal suspensions which eventually results in spontaneous precipitation, a process which occurs not only in chemically well-defined colloids, but is often observed in sera. Bechold has referred to this as "das Altern Kolloidaler Lösungen."

Of great interest, furthermore, in connection with the physical properties of complement is the discovery made by Jacoby and Schütze⁷⁰ that complement can be inactivated by shaking. This astonishing observation has been confirmed by Zeissler,⁷¹ Noguchi and Bronfenbrenner,⁷² Ritz,⁷³ and others. It appears, according to these observers, that guinea pig serum, when subjected to active shaking, can eventually be robbed thereby of its activating properties. The success of such experiments depends somewhat upon the concentration of the serum, and is best observed in a dilution of 1 part to 10 parts of salt solution. Under such conditions complete inactivation may be observed within 20 to 25 minutes. Between the inactivation of complement by heat and that which results from

⁶⁸ Gramenitski. *Loc. cit.*, p. 504.

⁶⁹ Traube. *Zeitschr. f. Imm.*, Vol. 9, 1911, and *Biochem. Zeitschr.*, 1908.

⁷⁰ Jacoby and Schütze. *Zeitschr. f. Imm.*, Vol. 4, 1910.

⁷¹ Zeissler. *Berl. kl. Woch.*, No. 52, 1909.

⁷² Noguchi and Bronfenbrenner. *Journ. of Exp. Med.*, Vol. 13, 1911.

⁷³ Ritz. *Zeitschr. f. Imm.*, Vol. 15, 1912.

shaking, there are certain similarities which seem to strengthen the opinion regarding the nature of heat inactivation which we have cited above. For it has been variously shown that prolonged shaking of protein solutions, like heating, gradually leads to coagulation. It would be important to determine whether or not the inactivation by shaking, like that produced by heat, is accompanied by a fall of surface tension.

ALEXIN OR COMPLEMENT FIXATION

The controversy regarding the multiplicity of alexin and the existence of a "complementophile group" cannot, of course, be regarded as closed, however much we may lean toward the acceptance, of Bordet's point of view, since German experimenters of eminence still adhere to the Ehrlich interpretations. Moreover, it is, of course, extremely difficult to disprove such an assumption as that of the "polyceptor" conception of the complementophile group. However, we may safely assert that the functional unity of complement (and, after all, that is all that Bordet has maintained) is being upheld by the constantly increasing evidence in its favor which is being furnished by the practical and experimental application of the phenomenon of "alexin fixation" described, in 1901, by Bordet and Gengou.⁷⁴ It will be well to bear in mind that this phenomenon should be strictly distinguished from the so-called "complement deviation" ("Ablenkung"), described by Neisser and Wechsberg. The latter was advanced as an explanation of the inactivity of bactericidal sera when used in too great concentration, as described in another place (p. 160) (Neisser and Wechsberg phenomenon), and has been variously utilized as support for the assertion that alexin can unite with unattached sensitizer. It is regarded by most observers, moreover, as untenable in the light of later investigation. In spite of this, the term "Komplement-Ablenkung" has been employed by a number of German writers (see Citron, Vol. 2, "Kraus und Levaditi Handbuch") as synonymous with "fixation" in the sense of Bordet and Gengou.

The phenomenon of Bordet and Gengou, briefly described, is nothing more than an experimental utilization of the fact which we have discussed at length, that alexin is fixed by antigen and antibody after union, but by neither alone.

The condition, as observed by them, may be best described by submitting the protocol of the first experiment detailed in their communication:

An emulsion of a 24-hour slant of plague bacilli was used as

⁷⁴ Bordet and Gengou. *Ann. de l'Inst. Past.*, 1901, Vol. 15, p. 289.

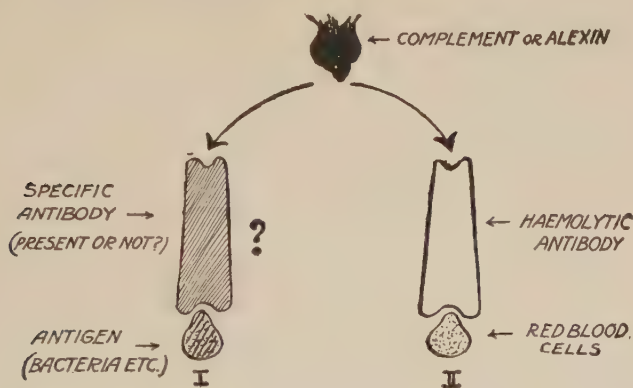
antigen, heated antiplague horse serum represented the antibody, and fresh guinea pig serum was used as alexin. A series of tubes was then prepared as follows:

1. Alexin + plague bacilli + inactivated antiplague serum.
2. Alexin + plague bacilli + inactivated normal horse serum.
3. Alexin + inactivated antiplague serum.
4. Alexin + inactivated normal horse serum.
5. Plague bacilli + inactivated antiplague serum.
6. Plague bacilli and normal horse serum.

These mixtures were left together for 5 hours and, at the end of this time, sensitized rabbit corpuscles were added to each tube. The result showed hemolysis in all the tubes except "1," in which there were plague bacilli, antiplague serum, and alexin, and in tubes 5 and 6, which had contained no alexin from the beginning.⁷⁵

It was plain, therefore, that the bacilli when specifically sensitized had become capable of absorbing alexin and preventing its subsequent action upon the sensitized erythrocytes. That the occurrence was not exceptional was shown by the fact that, in the same series, similar results were obtained with anthrax, typhoid, and proteus bacilli, and their respective antisera.

Schematized in accordance with the conceptions of Ehrlich our diagram would be as follows:



COMPLEMENT FIXATION SCHEMATIZED ACCORDING TO EHRLICH'S VIEWS.

If the antibody in I is present then complement is fixed by the antigen-antibody complex, and is no longer free to act upon the hemolytic complex II. In the same way antigen I could be determined if a known antibody I were used. For, in the absence of either of these parts of the complex I complement would remain unfixed and free to act on complex II.

We represent the phenomenon graphically in the symbols of Ehrlich merely because they facilitate clearness of exposition.

In the presence of both parts of Complex I the alexin is held and

⁷⁵ We will see later that unsensitized bacteria in emulsion will non-specifically fix small amounts of complement.

is no longer available for Complex II. If either of the reacting parts, antigen or sensitizer, of Complex I are lacking the alexin is left unfixed and free to react with Complex II.

With this technique Bordet and Gengou were able to demonstrate, by indirect experiment, the presence of specific sensitizers in the sera of animals immunized with various bacteria, a fact which was, of course, surmised but had been amenable to proof heretofore only in the case of bacteria like the spirillum of cholera in which lysis under the influence of immune serum and alexin could be directly observed under the microscope. The practical possibilities of their method were, of course, immediately apparent. By the use of a known antigen specific sensitizers can be demonstrated in this way, and, vice versa, in the presence of a known antibody, the method will serve to identify the nature of a doubtful antigen. Thus bacterial differentiation can be carried out by adding to the suspected bacteria, in emulsion, a small quantity of a known antiserum and alexin, and determining whether or not the alexin has become fixed. And, conversely, Bordet and Gengou⁷⁶ have more recently utilized the method in support of their claim of the specific etiological importance of the bacillus isolated by them from whooping cough, by showing that the serum of children suffering from this disease formed a specific alexin-fixing complex when treated with the bacillus.

The phenomenon of Bordet and Gengou thus found rapid practical application in the diagnosis of a number of infectious diseases, and has, of course, attained great clinical importance in the diagnosis of syphilis in the form of the "Wassermann" reaction and its many modifications. Before discussing these practical features in greater detail, however, it will be useful to discuss more particularly the many important theoretical considerations which have followed in the train of the complement-fixation phenomena.

A year after the publication of Bordet and Gengou's paper Gengou⁷⁷ made another fundamentally important observation by showing that complement or alexin fixation was not limited to the complexes of cellular antigens and their antibodies, but that the sera of animals immunized with dissolved proteins (animal sera, etc.), when brought together with their specific antigens, likewise formed combinations which fixed alexin. Thus egg-white or dog serum, brought together with "anti-egg-white" or "anti-dog" rabbit serum, respectively, strongly fixed alexin, whereas neither the antigenic substances nor the antisera exerted such fixation alone. The interpretation put upon this by Gengou was the following: "In sera obtained by injecting rabbits with large doses of cow's milk, etc., there are, in addition to the precipitins of Bordet and Tschistovitch, substances

⁷⁶ Bordet and Gengou. *Ann. de l'Inst. Past.*, 1906.

⁷⁷ Gengou. *Ann. de l'Inst. Past.*, 16, 1902.

analogous to the sensitizers described by Bordet in bacteriolytic and hemolytic sera, and later found in the majority of antimicrobial sera." The important point in this interpretation is that Gengou conceived the existence of antiprotein sensitizers, in addition to the precipitins, formed as a response to immunization with amorphous protein. Moreschi⁷⁸ soon confirmed Gengou's experimental determinations, and Neisser and Sachs⁷⁹ took the further logical step of applying this knowledge to the determination of proteins for forensic purposes. This, too, we will further discuss when we speak of the practical features of these phenomena. It thus appears that the fixation of alexin is a generalized property of all mixtures in which an antigen is brought into contact with its specific antibody, whether the antigen is in the form of the whole bacterial or other cell, or in that of a dissolved protein, animal serum, or egg-white, etc.

The observation of Gengou, though for a time insufficiently valued, has had a profound influence upon the subsequent understanding of serum reactions. The fundamental importance of this work was not fully recognized until his studies had found logical continuation in the investigations of Gay⁸⁰ and in those of Moreschi.

Moreschi⁸¹ studied the antihemolytic properties possessed by the serum of a rabbit which had been treated with normal goat serum. He found that such a serum had distinct anticomplementary powers when it was added to a hemolytic system of ox blood sensitizer (obtained against ox blood from rabbits), and goat complement. With such a hemolytic system, however, there was anticomplementary action only against goat complement and not against rabbit or guinea pig complement. If, however, he used a hemolytic system in which the amboceptor or hemolytic sensitizer employed was one obtained from a goat, the serum was anticomplementary for all complements which were used. Moreschi concluded from this that the apparent anticomplementary action of the serum could not be interpreted as the action of a specific anticomplement in the sense of Ehrlich, but that it resulted from the reaction which took place as the consequence of union of the antibody in the anti-goat rabbit serum and goat protein, which was introduced into the tubes, in the first case with the complement, and in the second with the amboceptor. He proved his contention by obtaining similar universal anticomplementary action when he added a little normal goat serum to the tubes set up as above described. It is plain, therefore, that anticomplementary action can be explained in observed cases by the simple consideration of the phenomenon of Gengou. Similar findings were later recorded by Muir

⁷⁸ Moreschi. *Berl. kl. Woch.*, No. 37, 1905.

⁷⁹ Neisser and Sachs. *Berl. kl. Woch.*, No. 44, 1905.

⁸⁰ Gay. *Centralbl. f. Bakt. I Orig.* Vol. 93, 1905, p. 603.

⁸¹ Moreschi. *Berl. kl. Woch.*, 1905, No. 37, *ibid.*, No. 4, 1906.

and Martin,⁸² and it may well be doubted, as a result of these and other researches, whether we are at all justified in assuming the existence of anticomplements.

The work of Gay, published independently in the same year as that of Moreschi, has, in a general way, the same significance, but Gay recognized the relation of the conditions observed by him to the precipitin reaction, a feature absent from both the original study of Gengou and the work of Moreschi. Gay noticed that an inactivated hemolytic immune serum, left for some time in contact with its specific cells, and then separated from them by centrifugation, would often possess anticomplementary or anti-alexin properties. He further noted that after such a serum had been freed from the cells by a short centrifugation, if it was again vigorously centrifugalized, a slight, cloudy sediment would appear at the bottom of the tubes. If this sediment was removed the serum lost its alexin-fixing properties. He recognized that the precipitate formed in these tubes was a specific precipitate resulting from the union of a precipitinogen and its antibody. The reaction was due entirely to the fact that insufficient washing of the cells used in producing the hemolysis, gave rise to the formation of precipitin against the serum of the animal from which the cells had been taken, and subsequently insufficient washing of the cells of this same species employed in the tests furnished enough antigen to give a precipitin reaction in the tubes in which the inactivated hemolytic (and precipitating) serum was mixed with the cells. Subsequently, numerous investigations⁸³ have shown Gay's interpretation to be correct, and we may now accept it as a fact that precipitates formed by the union of specific antigen with its antibody possess the power of fixing alexin and that, in a general way, this fixation is proportionate in energy to the amount of precipitate which is formed.

Gay utilized his results primarily to contradict certain assertions of Pfeiffer and Friedberger concerning antibacteriolytic substances supposed to occur in normal sera. These authors had found that, if normal sera possessing no "antagonistic" properties in the first place were left in contact with certain bacteria, they acquired antibacteriolytic properties for these particular bacteria. Thus normal inactive rabbit serum, left in contact with typhoid bacilli, and again separated from the bacteria, now prevented the lysis of sensitized typhoid bacilli if tested by the intraperitoneal method spoken of as the Pfeiffer reaction. Sachs⁸⁴ applied these observations to analogous hemolytic reactions and obtained similar results. He found that, if normal, inactive rabbit serum was left in contact with sheep

⁸² Muir and Martin. *Journ. of Hyg.*, Vol. 6, 1906. See also Muir's "Studies on Immunity," Froude, London, 1909.

⁸³ Dean. *Zeitschr. f. Imm.*, I, Vol. 13, 1912.

⁸⁴ Sachs. *Deut. med. Woch.*, 1905, No. 18.

or guinea pig corpuscles, it acquired the property of preventing the hemolysis of these corpuscles if, later, it was brought together with them in the presence of specific hemolysin and alexin. Gay now showed by experiment that Sachs' method was referable to insufficient washing of the corpuscles. When, in the first contact, the rabbit serum was exposed to the sheep corpuscles, a certain amount of sheep serum adherent to the cells was carried over into the rabbit serum. This sheep antigen later reacted with the antish sheep precipitin present in the hemolytic immune serum and, in this way, fixed alexin and prevented hemolysis.

It seems that the analysis of Gay is correct, and that Sachs' conclusion as well as those of Pfeiffer and Friedberger, by analogy, cannot be taken as demonstrating the existence of specific anticomplements or anti-amboceptors. Gay has further offered the same mechanism as an explanation of the Neisser-Wechsberg phenomenon, which has been discussed in another place.

To summarize, then, we have learned that there are a number of varieties of specific alexin absorption or fixation processes, one that is exerted by cells treated with specific sensitizer, be they blood or bacterial, the other that which occurs when unformed protein is brought into contact with its specific antiserum. The latter has been correlated with the precipitin reaction, in that it has been found that, whenever a specific precipitate is formed in such reactions, it is this precipitate on which the fixation depends. On the other hand, it is necessary to note that the formation of a precipitate is by no means necessary for the fixation, for, as is well known, if a series of precipitin tubes are set up, in each successive one of which the amount of antigen is diminished, a degree of dilution will soon be reached at which no visible precipitate will occur, but which nevertheless will show alexin fixation. The following is an illustration of such an experiment:

| Sheep serum + antish sheep serum | Precipitate | Fixation of 0.5 c. c. guinea pig complement |
|-------------------------------------|-------------|---|
| 0.5 c. c. (1:20) + 0.5 c. c. | + | Complete |
| 0.5 c. c. (1:50) 0.5 c. c. | ++ | Complete |
| 0.5 c. c. (1:100) 0.5 c. c. | +++ | Complete |
| 0.5 c. c. (1:200) 0.5 c. c. | +++ | Complete |
| 0.5 c. c. (1:500) 0.5 c. c. | +++ | Complete |
| 0.5 c. c. (1:1,000) 0.5 c. c. | + | Complete |
| 0.5 c. c. (1:2,000) 0.5 c. c. | + | Complete |
| 0.5 c. c. (1:5,000) 0.5 c. c. | | Partial |
| 0.5 c. c. (1:10,000) 0.5 c. c. | | Partial |
| 0.5 c. c. (1:20,000) 0.5 c. c. | | None |

From such experiments it follows moreover that the fixation of alexin, carefully titrated, is a more delicate method of determining

the presence of an antigen or, vice versa, of an antibody than is the observation of a visible precipitate, a fact which has been made use of, as we have mentioned, by Neisser and Sachs and others for forensic antigen determinations.

It should also be remembered that, if to such a precipitate there is added an excess of the antigen, the precipitate may be partially dissolved, and this dissolved precipitate, as Gay⁸⁵ has shown, may possess fixation properties. This, too, accounts for the fact, observed by a number of workers, that if, in a series of precipitin tests the supernatant fluids and the washed precipitates are separately examined for alexin fixation, the fixation properties reside entirely in the precipitates except in those tubes in which a considerable excess of antigen was used and in which, as in tubes 1 and 2 of the preceding protocol, the precipitates were relatively slight. The subject, though involved, is worthy of detailed consideration in this place since it seems to us to have an important bearing on certain theoretical conceptions which will be taken up below.

The important question now arises: what is the nature of the alexin fixation by the complexes formed by unformed proteins with their antibodies and, more especially, what is the nature of the alexin fixation exerted by specific precipitates? There have been much experimentation and speculation concerning this, and a number of different views are held. Gengou assumed, as we have seen, that this fixation, as studied by him, was entirely analogous to the fixation by sensitized bacterial or blood cells. He expressed the belief that treatment of an animal with an unformed protein produced not only specific precipitins but also specific sensitizers, analogous to those produced in response to treatment with bacterial or other cells. He noticed the parallelism between the quantity of the precipitate formed and the alexin fixation, but did not associate the two processes.

His conception of specific antiprotein sensitizers was accepted by a number of workers, and Wassermann and Bruck,⁸⁶ Friedberger⁸⁷ and several others brought out the facts that actual precipitate formation is not a necessary criterion of fixation. Thus the last-named writer showed that the precipitating power of a serum may be destroyed by moderate heat without a corresponding destruction of its fixing property. A similar independence of the precipitation from the complement-fixing property, in the presence of an antigen, has been observed by Muir and Martin.⁸⁸

⁸⁵ Gay. *Univ. of Cal. Public. in Pathol.*, Vol. 2, No. 1, 1911.

⁸⁶ Wassermann and Bruck. *Mediz. Kl.*, 1905, Vol. 1, No. 55.

⁸⁷ Friedberger. *Deut. med. Woch.*, 1906, No. 15.

⁸⁸ Muir and Martin. *Jour. of Hyg.*, Vol. 6, 1906.

Gay,^{89 90} also, though he was the first definitely to associate precipitin formation with the alexin-fixing property and, indeed, determined a rough parallelism between the amount of precipitate and the degree of alexin fixation, has nevertheless recently declared himself in favor of the assumption of the presence in protein antisera of two antibodies, the alexin-fixing lysins and the precipitins. This he does on the basis of certain experiments from which he concludes that the antigen-antibody complex which fixes alexin is distinct from the precipitin-precipitinogen complex, but is usually "brought down in its formation in such a way as to simulate fixation by the precipitate." Nicolle⁹¹ goes even further than this in declaring that the "coagulins" or precipitins are "anticorps bons," which prevent the action of the albuminolysin upon the antigen, thereby inhibiting the liberation of poisonous cleavage products.

It seems to the writer^{92 93} that the assumption of a separation between the precipitin and the albuminolysin is a needlessly complicated interpretation of the phenomena. In order to elucidate this point a comparison was made between the fixing properties of a mixture of a protein (sheep serum) and its antibody and a mixture of typhoid filtrate and antityphoid serum in which it is known that both precipitins and antibacterial sensitizers are present. It was shown that, as stated before, in the former mixture the alexin-fixing property resided entirely in the precipitate, whereas in the latter case both the precipitate and the supernatant fluid fixed alexin. From this it seems to follow that immunization with the more complex cellular elements has given rise to the precipitating antibody present also in the antsheep serum, and, in addition to this, to sensitizers which are not precipitable (remaining in the supernatant liquid) and not present in the antsheep serum. The precipitates, moreover, were found to fix "end-piece" and "mid-piece," fractions of alexin, in the same way as these are fixed by sensitized cells.

Without going into further complicated detail, it would seem to us⁹⁴ to be justified that we look upon the so-called precipitins not as separate antibodies but as identical with so-called albuminolysins. They unite with the antigen, producing an alexin-fixing complex. Since both reacting bodies are colloidal in nature, they precipitate each other in the test tube, but, following the laws governing other mutually precipitating colloids, they do so only when brought to-

⁸⁹ Gay. *Loc. cit.*

⁹⁰ Also *Univ. of Cal. Publ. in Pathol.*, Vol. 2, No. 1, 1911.

⁹¹ Nicolle. Ref. in *Bull. de l'Inst. Past.*, Vol. 5, 1907.

⁹² Zinsser. *Journ. Exp. Med.*, Vol. 15, 1912.

⁹³ Zinsser. *Proc. of Soc. of Exp. Biol. and Med.*, April, 1913.

⁹⁴ Zinsser. *Journ. of Exp. Med.*, Sept., 1913.

gether in concentrations which lie within definite zones of relative proportions. The visible precipitation would seem, therefore, to be merely a secondary phenomenon, the essential one being the union of an antigen with a sensitizer by which it is rendered amenable to the action of the alexin. This would enable us to comprehend also the experiments of Friedberger, discussed in the section on anaphylaxis, in which it was shown that the action of alexin upon precipitates gives rise to the formation of toxic bodies just as this occurs when alexin acts upon sensitized cells. It leads, moreover, to a comprehension of the processes of the digestion of intravascularly introduced foreign proteins, which are rendered amenable to the digestive action of the alexin by the antibodies spoken of as precipitins, which functionally and in structure are conceived as identical with other sensitizers.

Dean,⁹⁵ who has lately analyzed the relation between precipitation and alexin fixation on the basis of extensive experimentation, comes to the conclusion that the proportions of antigen and antibody which are favorable for rapid and complete precipitation do not favor the most complete alexin fixation. He states that the two reactions do not run a parallel course but believes that this does not mean that they are necessarily distinct phenomena. He says: "They represent two phases of the same reaction . . . a flocculent precipitate represents the final stage of a change which can be recognized in its earliest and incomplete stage by means of a complement fixation."

Our view differs from this only in that we believe that the precipitation is merely a secondary, colloidal phenomenon, which may, or may not, coincide with the phase of greatest alexin fixation, according to other fortuitous conditions which may favor or retard flocculation. Indeed, if our view be accepted, rapid compact precipitation may possibly be assumed to interfere with alexin fixation in that it would inhibit perfect contact of the alexin with the antigen-antibody complexes.

Another view of the mechanism of alexin fixation is that which has been advanced by Neufeld and Haendel.⁹⁶ These workers have found that sensitized cholera spirilla will fix hemolytic complement at 0° C., whereas the same bacteria at 37° C. will fix both the hemolytic and the bactericidal complement. They conclude from this that the fixation at 37° C. was brought about by virtue of the bactericidal amboceptor, whereas at 0° C. fixation was brought about by an antibody which is distinct from amboceptor or sensitizer. They believe from this and other observations, which we cannot consider in detail, that alexin fixation may be brought about by a special fixing anti-

⁹⁵ Dean. *Zeitschr. f. Imm.*, Vol. 13, 1912.

⁹⁶ Neufeld and Haendel. *Arb. a. d. kais. Gesund.*, Vol. 28, 1908.

body, the "Bordetscher Antikörper," which is not identical with any of the other known antibodies.

In all experiments which deal with alexin fixation by specific antigen-antibody complexes it is of the greatest importance that we should guard against the errors easily introduced by fortuitous non-specific antihemolytic agencies. Thus there are a number of factors which will interfere with the functionation of alexin upon a sensitized antigen, either by direct non-specific absorption of the alexin itself or by producing physical conditions in the presence of which alexin cannot act.

Thus many animal tissue cells, in emulsion, will absorb alexin, and the same property may be possessed by tissue extracts. Von Dungern⁹⁷ was the first to call attention to this, and his observations have been variously confirmed. Muir⁹⁸ showed that the stromata of hemolyzed red blood cells exert strong anticomplementary action, and that this is due to a firm union with the complement. It is not unlikely that the action of cells in this respect is referable to their lipoidal contents. This suggestion was first made by Landsteiner and von Eisler,⁹⁹ who found that the petroleum-ether extracts of red blood cells possessed strong anticomplementary action which, to a limited extent, was specific toward the particular corpuscles from which the extracts had been made. Similar observations have been made by Noguchi,¹⁰⁰ who speaks of the substance he extracts as "protectin." In general, the protective action of the lipoidal extracts seems to depend largely upon cholesterin, and, since this substance is present to some extent in many tissues, their antihemolytic action is easily understood. In another section we have discussed the similar neutralizing action of lipoidal substances upon poisons of various kinds (saponin, tetanolyisin, and snake poison), but, as we have noted there, the neutralizing properties of the extracts do not, as a rule, equal those of the whole tissues.¹⁰¹ It is not unlikely that in such cases as Landsteiner suggests the potent agent is not the lipid itself but rather a lipid-protein combination, a class of substances of which we know very little, but the importance of which, in many phases of serum reactions, seems assured.

We have already mentioned that yeast cells may absorb alexin. And it has been found by Wilde¹⁰² and others that almost all bacteria in emulsion may possess varying degrees of alexin-fixing properties even though unsensitized. There seems to be no regularity either qualitatively or quantitatively in regard to this, but the fixa-

⁹⁷ Von Dungern. *Münch. med. Woch.*, Nos. 20 and 28, 1900.

⁹⁸ Muir. "Studies in Immunity," London, Vol. 19.

⁹⁹ Landsteiner and von Eisler. *Wien. kl. Woch.*, No. 24, 1904.

¹⁰⁰ Noguchi. *Journ. Exp. Med.*, Vol. 8, 1906, p. 726.

¹⁰¹ See also Ivar Bang, "Biochemie der Lipide."

¹⁰² Wilde. *Berl. kl. Woch.*, 1901, Vol. 38, and *Archiv f. Hyg.*, 39, 1902.

tion is usually sufficiently marked to render the use of whole bacteria unreliable for specific fixation experiments. For this reason, as we will see, bacterial extracts must be used in such work unless careful quantitative controls are made. Upon what this fixation depends it is difficult to determine. It may be that it is purely non-specific and due to absorption of the fine emulsion of the bacteria comparable to that observed on the part of kaolin or quartz sand emulsions, or, possibly fixation by such bacterial emulsions may occur because of the small amounts of normal sensitizer almost always present in the serum employed as alexin.

Apart from the lipoids, a number of other substances have been found to fix alexin and exert consequent antihemolytic action. Thus Landsteiner and Stankovic,¹⁰³ and Landsteiner and von Eisler¹⁰⁴ describe the anti-alexin action of various proteins coagulated or precipitated. They refer this action not to particular chemical structure but to the colloidal state, since they obtained similar antilytic action with such inorganic emulsions as quartz sand and kaolin (aluminium-orthosilicate). Since anticomplementary action has, moreover, been noted in the case of a large number of extracts of such materials as wool, leather, etc., it is clear that the methods of alexin fixation, as applied to the forensic differentiation of blood, must be carefully controlled with this point in view.¹⁰⁵

Among the most practically important non-specific agencies which fix alexin there are some which appear under certain conditions in normal serum. Noguchi¹⁰⁶ has found that serum will often develop anticomplementary properties as a consequence of heating during the process of inactivation. On more detailed investigation he determined that the anticomplementary action increased as the serum was heated to about 90° C. Above this temperature it is destroyed. He refers this property to the serum lipoids, since he was able to remove it by extraction with ether, the ether extract possessing the same anticomplementary power as the original serum.

Neisser and Döring¹⁰⁷ have noticed anti-alexin or anticomplementary properties of human sera which were destroyed on heating, and which they associate with disease of the kidneys, since they noted it in sera of uremic patients. Browning and McKenzie¹⁰⁸ have observed a similar heat-sensitive anti-alexin action on the part of normal serum, and the subject has been studied by Zinsser and Johnston.¹⁰⁹ It was found that all normal sera will develop anti-

¹⁰³ Landsteiner and Stankovic. *Centralbl. f. Bakt.*, 1906, Vols. 41 and 42.

¹⁰⁴ Landsteiner and von Eisler. *Wien. kl. Woch.*, 1904, No. 24.

¹⁰⁵ Uhlenhuth. *Deut. med. Woch.*, 1906, Nos. 31 and 51, and *Centralbl. f. Bakt.*, 1906, I, Ref., Vol. 38.

¹⁰⁶ Noguchi. *Journ. of Exp. Med.*, Vol. 8, 1906, p. 726.

¹⁰⁷ Neisser and Döring. *Berl. kl. Woch.*, 1901, No. 22.

¹⁰⁸ Browning and McKenzie. *Journ. of Path. and Bact.*, Vol. 13, 1909.

¹⁰⁹ Zinsser and Johnston. *Journ. of Exp. Med.*, Vol. 13, 1911.

alexie properties on preservation at room temperature within a few days, and more slowly but no less regularly in the ice chest. This anti-alexin is destroyed on heating to 56° C., and may be precipitated out with the globulins of the serum. There appeared in these studies no particular association between the anti-alexie property and nephritis.

The action of alexin upon sensitized cells may be prevented, also, by physical or chemical conditions without actual fixation or binding of the alexin. We refer to the effects of the addition of salts, problems which have been considered above.

CHAPTER VIII

PRACTICAL APPLICATIONS OF THE COMPLEMENT-FIXATION METHOD

THE WASSERMANN REACTION

THE principle of specific alexin fixation has been practically utilized in the diagnosis of disease and in the forensic determination of the nature of spots of blood or other protein material.

Soon after Bordet and Gengou's experiments Wassermann and Bruck¹ showed that bacterial extracts could be successfully substituted for whole bacteria in these reactions. Citron,² too, made similar observations, and, indeed, we now know that the use of bacterial extracts is more suitable for these experiments than are emulsions of whole bacteria, since, as we have mentioned above, bacterial emulsions may often fix small amounts of complement of themselves (without specific sensitization), thereby confusing the results of the reaction.

On the basis of their experience with bacterial extracts Wassermann and Bruck³ then determined that complement fixation could be carried out in tuberculosis when the various tuberculin preparations were used as antigen.⁴ These investigations fell into the period during which active research upon the *Spirochæta pallida* in syphilis was going on, and it occurred to Wassermann that the technique of complement or alexin fixation might be utilized in the diagnosis of syphilis. Together with Neisser and Bruck⁵ he subjected this idea to experimental test. The publication of their first results appeared in 1906. They used in their experiments the syphilitic monkeys which were being observed in Neisser's clinic. Their method consisted in mixing inactivated serum from syphilis-inoculated monkeys with organ extracts, serum, etc., of syphilitic human beings, and

¹ Wassermann and Bruck. *Med. Klinik*, Vol. 55, 1905.

² Citron. *Centralbl. f. Bakt.*, Vol. 41, 1906.

³ Wassermann and Bruck. *Deut. med. Woch.*, No. 12, 1906.

⁴ Complement fixation in tuberculosis is not yet on a practical or reliable basis. Recent claims of Besredka (*Ann. Past.*, 1913) for his new antigen promise a successful technique, but no extensive confirmation has followed up to the present time.

⁵ Wassermann, A. Neisser, and Bruck. *Deut. med. Woch.*, No. 19, 1906.

adding a small amount of fresh guinea pig complement. After these materials had been together for a certain time, sensitized red blood cells were added. If the complement was bound during the first exposure no hemolysis resulted and the reaction was regarded as positive. From their results they drew the following conclusions:

1. Immune serum from monkeys, produced by treatment with syphilitic material, will sensitize syphilitic material from human beings or monkeys, so that an alexin-fixing complex is formed.

2. Complement fixation results only when the syphilitic immune serum of monkeys is added to similar material from men or monkeys, but not when added to organ extracts of normal men or monkeys.

3. Normal monkey serum has no such action.

They concluded that their results justified them in assuming a specific fixation due to specific antisyphilitic immune bodies in the blood of the treated monkeys. They excluded experimentally the possibility of fixation by a precipitin reaction resulting from the treatment of the monkeys with human material. It might well have happened that precipitins against human protein appearing in the serum of the treated monkeys might subsequently react with the human protein material used as antigen, a complement-fixing complex resulting. This, however, was excluded by the fact that they obtained positive reactions only when the human material was obtained from luetic lesions.

The same authors, with Schucht,⁶ very soon after this, extended their method to the diagnosis of syphilis in human beings. The same thing had been done shortly before their publication appeared by Detre⁷ on a smaller material. By these and many other investigations it was very soon shown that syphilis may be reliably diagnosed by complement fixation when extracts of the syphilitic organs, employed as antigen, are mixed with the inactivated serum of syphilitic individuals. It was incidentally shown by Wassermann and Plaut⁸ that the reaction could be obtained not only with blood serum but also with spinal fluid in paralytic cases.

It was generally assumed, at this time, that the reaction in syphilis depended, as in the case of other infections, upon the presence in the syphilitic serum of specific antibodies. For it seemed reasonable to suppose that the specific antigen obtained in the extracts was derived from the extraction of large numbers of spirochetes demonstrable in the extracted organs.

This, of course, is the most logical and simple theoretical conception of the reaction, and is justified on the basis of analogy. Un-

⁶ Wassermann, Neisser, Bruck, and Schucht. *Zeitschr. f. Hyg.*, Vol. 55, 1906.

⁷ Detre. *Wien. kl. Woch.*, Vol. 19, No. 21, 1906.

⁸ Wassermann and Plaut. *Deut. med. Woch.*, No. 44, 1906.

fortunately, however, it was soon found by a number of workers, Marie and Levaditi,⁹ Weygant, Kraus and Volk, Landsteiner, Müller, and Pötzl,¹⁰ and others that antigens perfectly capable of fixing complement in the presence of syphilitic serum could be produced from normal organs.¹¹

Theoretically it must be admitted that we are very much in the dark at present. The fact, now entirely unquestionable, that the sera of syphilitic patients will give fixation with antigens derived from extracts of normal organs, as well as from those of syphilitic organs, seems to throw doubt upon the simple specific antigen-antibody conception at first held.

In order to understand the questions involved in the theories of the Wassermann reaction as at present conceived it will be necessary to consider the types of antigen which are now employed.

Wassermann's original method of antigen preparation consisted in using the liver or spleen of a congenitally syphilitic fetus. The organs were finely divided and emulsified in 4 to 6 parts of normal salt solution. This mixture was shaken for 24 hours, centrifugalized, and the clear supernatant fluid used as antigen. Later the specific organ substances were extracted by Porges and Meier¹² in five times the volume of absolute alcohol for 24 hours. This alcoholic extract was evaporated *in vacuo* and the residue taken up in salt solution and shaken until an even suspension resulted.

After it had been discovered that normal organ extracts could serve as antigen as well as the extracts of syphilitic organs, Landsteiner, Porges and Meier, and others, introduced antigens produced by alcoholic extraction of normal organs of animals and of man. Landsteiner introduced the alcoholic extract of normal guinea pig organs, especially extracts of the heart and liver, and Weil and Braun¹³ made use of extracts of normal human organs. There are various methods of preparing extracts for this purpose. We may mention, to illustrate these methods, the one suggested, first, we believe, by Noguchi, a procedure which is applicable to the extraction of normal human organs (spleen), beef hearts, and guinea pig hearts. The finely divided or triturated organ substance is shaken up with five times its weight of absolute alcohol and allowed to stand in the

⁹ Marie and Levaditi. Cited from McIntosh and Fildes' "Syphilis." Longmans & Co., 1911, p. 94.

¹⁰ Landsteiner, Müller, and Pötzl. *Wien. kl. Woch.*, Vol. 20, 1907.

¹¹ An extensive historical review of the development of the Wassermann reaction is found in the book of Boas, "Die Wassermannsche Reaktion," Karger, Berlin, 1911. Since these earlier publications have appeared the literature of the Wassermann reaction has become very extensive. It is enumerated more fully than we can afford space for here in the book of Noguchi ("Serum Diagnosis of Syphilis") and that of Boas, mentioned above.

¹² Porges and Meier. *Berl. kl. Woch.*, No. 15, 1908.

¹³ Weil and Braun. *Berl. kl. Woch.*, No. 49, 1907.

incubator at 37.5° C., for from 5 to 7 days. At the end of this time it is filtered through cheesecloth and then through coarse paper, and the filtrate placed in a large crystallizing dish in which it is evaporated at room temperature with the aid of an electric fan. A gummy yellow residue is left, which is then taken up in as small a quantity of ether as possible. This ether solution is then precipitated with 4 times its volume of acetone, in consequence of which there is a profuse precipitation of coarse white flakes. This acetone-insoluble substance, which is at first white, later yellowish, in color, is the stock antigen. A little of this is taken up in a very small quantity of ether, and this ethereal solution is shaken up in salt solution until the ether has evaporated or the material has gone into very fine colloidal suspension in the salt solution. This is the antigen ready to be used.

It is immediately evident that these antigenic substances must consist very largely of lipoidal extractives of the organ substances, and it has been found that such antigen contains sodium oleate, lecithin and cholesterin. Indeed, Porges and Meier have claimed that a 1 per cent. solution of commercial lecithin may be used with success. Browning and Cruikshank¹⁴ have found further that the addition of small amounts of cholesterin to syphilitic antigen very largely increases its specifically diagnostic value, and this idea has since been utilized more especially by Sachs,¹⁵ Walker and Swift,¹⁶ and others. Sachs, especially, has obtained excellent antigens in the following way: 1 gram of moist guinea pig heart substance was extracted with 5 c. c. of alcohol and left at room temperature for twelve hours or in the ice box for two days; it was then filtered and 0.5 to 1 per cent. of cholesterin was added; frequently the alcohol extract had to be diluted two or three times before use. Sachs and Rondoni¹⁷ have also recommended artificial mixtures of lipoids containing sodium oleate, lecithin, and oleic acid.

The fact that cholesterin added to alcoholic organ extracts increases the antigenic value of these for the Wassermann reaction is all the more curious inasmuch as cholesterin alone has practically no antigenic action. Walker and Swift have recommended an antigen in which alcoholic extracts of human or guinea pig hearts were made up to 0.4 per cent. of cholesterin, 0.4 per cent. having been found by comparative test to be the most favorable concentration. Cholesterin-liver extracts or even alcoholic extracts of syphilitic livers without cholesterin were found to be inferior in specific antigenic value to 0.4 per cent. cholesterin-heart antigens. From the experience of many investigators it now seems unquestionable that additions of

¹⁴ Browning and Cruikshank. *Journ. of Path. and Bact.*, Vol. 16, 1911.

¹⁵ Sachs. *Berl. kl. Woch.*, No. 46, 1911.

¹⁶ Walker and Swift. *Journ. of Exp. Med.*, Vol. 18, 1913.

¹⁷ Sachs and Rondoni. *Zeitschr. f. Imm.*, Vol. 1, 1909.

cholesterin increase the delicacy of the reaction in that more cases react positively with such an antigen than with the uncholesterinized preparations. The experience of Hopkins and Zimmermann, however, would indicate that great caution must be exercised when the reaction is done in this way, since occasional positive results are obtained with cases clinically not syphilitic. These workers believe that cholesterinized antigen is extremely useful, but advise its use only parallel with the ordinary lipoidal antigens and together with careful study of the clinical aspects of the case.

The fact that these antigens are non-specific in origin naturally necessitates careful determination of their usefulness before they are used. Before any antigen can be regarded as reliable, therefore, a titration must be carried out in the following way: Two series of tubes are prepared, in the first of which antigen and complement are added to normal serum, and in the second the same substances are added to known syphilitic serum. The antigen must, of course, be such that in no test tube does it cause alexin fixation in the presence of normal serum, but, in the quantities used, it must give fixation regularly with syphilitic serum. An example of such a titration may be tabulated as follows:

EXAMPLE OF ANTIGEN TITRATION

Antigen by Landsteiner's method: normal guinea pig heart freed from fat and ground up in a mortar. To each gram is added 5 c. c. of absolute ethyl alcohol and the mixture allowed to extract at 60° C. for 12 hours (or several days at 37.5° C.). It is then filtered through paper. The following titration is then carried out:

| A | Tube 1 | Tube 2 | Tube 3 | Tube 4 | Tube 5 |
|-------------------|--------|--------|--------|--------|--------|
| Normal serum..... | 0.2 | 0.2 | 0.2 | 0.2 | — |
| Antigen..... | 0.05 | 0.1 | 0.2 | 0.3 | 0.6 |
| Alexin..... | 0.1 | 0.1 | 0.1 | 0.1 | 0.1 |

| B | Tube 1 | Tube 2 | Tube 3 | Tube 4 | |
|-----------------------|--------|--------|--------|--------|--|
| Syphilitic serum..... | 0.2 | 0.2 | 0.2 | 0.2 | |
| Antigen..... | 0.05 | 0.1 | 0.2 | 0.3 | |
| Alexin..... | 0.1 | 0.1 | 0.1 | 0.1 | |

The volume in all of these tubes is brought to 3 c. c. with isotonic salt solution. After one hour at 37.5° C., sensitized red cells are

added to each tube.¹⁸ If the antigen is suitable in that it does not fix alexin by itself or in the presence of normal serum, hemolysis will result in all of the tubes of series A. If it is suitable in that it fixes in the presence of syphilitic serum, the tubes in series B will show no hemolysis; if there is slight hemolysis in B 1, it is inferred that 0.05 c. c. of the antigen is insufficient, and the smallest amount (0.1 c. c.), which completely fixes 0.1 c. c. of alexin in the presence of the positive serum, is the quantity used. Again the antigen may be able to cause hemolysis by itself if used in too large amounts. If this is the case in tube B 4, then this antigen is suitable only in amounts varying between 0.1 c. c. and 0.2 c. c.

The titration is done with varying quantities because too little antigen might fail in fixing the alexin, even if the serum were positively syphilitic, whereas too much antigen might possess alexin-fixing properties in itself, even in the presence of normal serum, or possibly without any serum at all, an attribute which is not uncommonly possessed by lipoidal extracts.

It is thus seen that Wassermann reactions can be carried out with antigens which do not contain extracts of syphilitic lesions or of the micro-organisms which give rise to syphilis. This fact alone would exclude the possibility of considering the fixation of complement as at present carried out in the Wassermann reaction as being due to a specific antigen-antibody union.

This conclusion is strengthened by the recent discovery that a specific antigen prepared from cultures of *Spirochaeta pallida* cannot be successfully used in diagnostic Wassermann tests. The first investigations of this kind were made by Schereschewsky,¹⁹ who used as antigen extracts of mixed cultures in which the spirochete was present; his results were inconclusive. Noguchi²⁰ later investigated this phase of the problem, preparing his antigens by the extraction of pure cultures and of syphilitic rabbit testicles in which the spirochetes were very profuse. He found that positive tests with such an antigen were obtained only in isolated cases of prolonged syphilis which had been thoroughly treated, and that the ordinary Wassermann reaction, as obtained in active cases, is not due to antibodies which combine specifically with the pallida antigen. Craig and Nichols²¹ also have found that cases of untreated syphilis which gave positive reactions with syphilitic liver extracts gave absolutely negative results when culture antigens were used.

¹⁸ Tube "5" is the antigen control which shows that the antigen in large amounts is neither anticomplementary nor hemolytic by itself. It is well, in addition, also to test out various amounts of the antigen and alexin, without either normal or syphilitic serum, to determine the largest amount of antigen which, by itself, is devoid of the actions mentioned above.

¹⁹ Schereschewsky. *Deut. med. Woch.*, 1909, p. 1653.

²⁰ Noguchi. *Journ. A. M. A.*, Vol. 58, 1912.

²¹ Craig and Nichols. *Journ. of Exp. Med.*, Vol. 16, 1912.

From these results also we may infer that the Wassermann reaction does not represent a fixation of alexin by the union of a specific syphilitic antigen with antibodies found against the *Spirochaeta pallida*. Noguchi concludes that it is caused by "lipotropic" substances in the sera of syphilitic human beings; a conclusion which is justified by the fact that the antigens used, all of them, contain large quantities of lipoids. It must be acknowledged, however, that we have no definite information concerning the nature of the reaction beyond this. Schmidt²² believes that it is a colloidal reaction, and depends upon the union of the serum globulins with the extract colloids in the antigen. In normal serum such a union is prevented by the albumins which act as a sort of protective colloid. In syphilitic serum the globulins are increased quantitatively or are changed qualitatively in the degree of their dispersion, or possibly in both characteristics. He regards the serum globulins in the Wassermann reaction as directly uniting with the extract colloid.

Levaditi and Yamanouchi²³ also conclude that the Wassermann reaction depends upon the union of two colloidal substances—one a non-proteid constituent of syphilitic serum (cholesterin derivatives or fatty acids), the other the lipoidal constituents of the antigen. Like others they found that the active substances in the antigenic extracts are non-protein and alcohol soluble.

It is interesting to note, moreover, that Porges and Meier²⁴ observed actual precipitation when syphilitic serum was added to lecithin emulsions. In consequence, attempts have been made to make the diagnosis of syphilis by direct precipitation of syphilitic serum by such emulsions of lecithin and of sodium glycocholate (Merck). The results of these investigations as well as those of Klausner,²⁵ who claims that syphilitic sera are more easily precipitated by distilled water than are normal sera, have led to no diagnostically reliable results, but they have seemed to show that the serum globulins are probably more plentiful and more easily precipitated out of syphilitic than out of normal sera.

The inference of many workers, therefore, has been that the Wassermann reaction is primarily due to the precipitation of (probably) globulin by the lipoidal colloids of the antigen, the resulting precipitate being capable of absorbing alexin. Jacobsthal²⁶ has examined mixtures of syphilitic serum and antigen by the ultramicroscopic method, and claims that precipitates are always present even when they are not macroscopically visible. Bergel,²⁷ who has re-

²² Schmidt. *Zeitschr. f. Hyg.*, Vol. 69, 1911.

²³ Levaditi and Yamanouchi. *C. R. de la Soc. de Biol.*, 1907, Vol. 63, p. 740.

²⁴ Porges and Meier. *Berl. kl. Woch.*, No. 15, 1908.

²⁵ Klausner. *Wien. kl. Woch.*, No. 7, 1908.

²⁶ Jacobsthal. *Münch. med. Woch.*, 1910.

²⁷ Bergel. *Zeitschr. f. Imm.*, Vol. 17, 1913.

ently suggested the importance of specific lipase production as a cause of hemolysis, suggests that the Wassermann reaction is due to fixation exerted by the products of the action of a specific lipase formed in the syphilitic body against "lues-lipoids." This theory is open to objections similar to those mentioned above, namely, that the antigen need not necessarily be a lues-lipoid, but may be derived from normal organs. Other theories have been brought forward by Bruck, Weil, Braun, Manwaring, and more recently by Rabinowitch.²⁸ The data supporting most of these theories are, as yet, too speculative to justify our discussion of them at any length. The only fact which seems established with any reasonable certainty is the independence of the Wassermann test from a specific antigen-antibody reaction in the usual sense.

Although the Wassermann reaction is thus apparently not based on those principles in the investigation of which it was discovered, its practical diagnostic value is not therefore diminished. For its proper performance any of the methods of antigen preparation considered above may be employed, provided that the usefulness of the preparation utilized is carefully controlled in each case as indicated. Since, of course, a hemolytic system is used in such tests as an indicator, it is necessary also to titrate sensitizer and alexin.

From what has been said in another place concerning the quantitative relations of alexin and amboceptor or sensitizer (see reference to work of Morgenroth and Sachs, p. 163), it is evident that the use of too strongly sensitized cells might result in hemolysis, if a slight fraction of alexin were left unbound by a weak syphilis reaction. Conversely the use of too large a quantity of alexin would result in hemolysis, since, even if the amount of syphilitic fixation were considerable, a sufficient excess of alexin might remain. The use of uniform amounts of fresh guinea pig serum in each case does not control this adequately, for different specimens of guinea pig serum may vary considerably in alexin content. In consequence, titrations of both sensitizer and alexin should be made. For practical purposes it is quite enough to titrate the hemolytic sensitizer every few weeks and use a stated amount in successive reactions. The alexin or complement can then be titrated individually for each set of reactions. Examples of such preliminary titrations follow:

Titration of Hemolytic Amboceptor or Sensitizer

Rabbit injected 3 times at 5-day intervals with washed sheep corpuscles, 3, 4, and 5 c. c., and bled 10 days after the last injection.²⁹

²⁸ Rabinowitch. *Centralbl. f. Bakt., Orig.*, 1914.

²⁹ In immunizing animals with blood cells for this or any other purpose it is necessary to wash the cells very carefully in salt solution. Unless this is

This serum is inactivated at 56° C. for 20 minutes.

| | Washed sheep corpuscles 5% emulsion in salt solution | Sensitizer | Fresh g. p. serum | Hemolysis |
|---|--|------------|-------------------------|-----------|
| 1 | 1 c. c..... | 0.01 | 0.1 | +++ |
| 2 | 1 c. c..... | 0.005 | 0.1 | +++ |
| 3 | 1 c. c..... | 0.003 | 0.1 | +++ |
| 4 | 1 c. c..... | 0.001 | 0.1 | +++ |
| 5 | 1 c. c..... | 0.0005 | 0.1 | ++ |
| 6 | 1 c. c..... | 0.0002 | 0.1 | ± |
| 7 | 1 c. c..... | | 0.1 | |
| 8 | 1 c. c..... | salt sol. | | |

In this case 0.001 c. c. still causes complete hemolysis of 1 c. c. of a 5 per cent. emulsion of sheep cells (volumetric measurement of cells sedimented in the centrifuge), and this amount ($\frac{1}{1000}$ c. c.) is called the "hemolytic unit" of sensitizer; two units are then used in the reactions.

Against these cells alexin can, in each case, be titrated as follows:

Alexin Titration:

Fresh Guinea Pig Serum Pipetted from Clot

| | Red cells 5% emulsion | Sensitizer as above determined | Guinea pig serum | Hemolysis |
|---|--------------------------|--------------------------------------|---------------------|-----------|
| 1 | 1 c. c. | 2 units (.002) | 0.1 c. c. | +++ |
| 2 | 1 c. c. | 2 units (.002) | 0.05 c. c. | +++ |
| 3 | 1 c. c. | 2 units (.002) | 0.025 c. c. | ± |
| 4 | 1 c. c. | 2 units (.002) | 0.01 c. c. | |

The smallest amount of alexin which completely hemolyzes the red cells (0.05 in this case) is the amount used. Since it is easier to measure larger volumes with accuracy, the alexin is diluted 1 to 10 in salt solution before use. A typical Wassermann reaction can then be carried out as follows:

done blood serum or plasma will be injected with them and the treated animal will respond by the formation not only of hemolysin but of precipitins for the serum proteins as well. When a subsequent hemolytic test is carried out, a precipitin reaction between the precipitin in the antiserum and serum adhering to the corpuscles will follow, and this, as we have seen, will fix alexin, obscuring other reactions which may be under observation.

SCHEME FOR WASSERMANN TEST

ADAPTED TO ORIGINAL WASSERMANN SYSTEM AFTER SCHEME OF NOGUCHI

| | Test with unknown serum | Test with known positive syphilitic serum | Test with known negative normal serum | Test without serum to control efficiency of hemolytic system |
|-----------------------------|---|---|--|---|
| Back row without antigen | Serum 2 c. c. + ● Complement .1 c. c. + Salt sol. 3 c. c. 2. | Total volume, 3 c. c. Serum .2 c. c. + ● Complement .1 c. c. + Salt sol. 3 c. c. 4. | Serum .2 c. c. + ● Complement .1 c. c. + Salt sol. 3 c. c. 6. | ● Complement .1 c. c. + Salt sol. 3 c. c. 8. |
| Front row with antigen | Serum .2 c. c. + ● Complement .1 c. c. + Antigen (required amount in 1 c. c. salt sol.) + Salt sol. 2 c. c. 1. | Total volume, 3 c. c. Serum .2 c. c. + ● Complement .1 c. c. + Antigen + Salt sol. 2 c. c. 3. | Serum .2 c. c. + ● Complement .1 c. c. + Antigen + Salt sol. 2 c. c. 5. | ● Complement .1 c. c. + Antigen + Salt sol. 2 c. c. 7. |

● = Test tube.

Place in water bath at 40° C. for one hour, then add to all tubes red blood cells and amboceptor. These are previously mixed so that 2 c. c. contains the equivalents of 1 c. c. of a 5 per cent. emulsion of sheep corpuscles and 2 units of amboceptor. Again expose to 40° C. If the serum tested is positive, tubes 1 and 3 should show no hemolysis, all the other tubes showing complete hemolysis in one hour.

Since many human sera normally contain small amounts of antish sheep sensitizer, it is the habit of many workers to add the sheep corpuscles, without the sensitizer or amboceptor, and incubate for a half-hour. If, at the end of this time, no hemolysis has occurred either in the front or the back row, then amboceptor may be added. This technique avoids the possible error introduced by an excess of amboceptor, a condition which easily occurs when any large amount is normally present in the serum and in addition to this 2 units are added as in the test described above.

The above represents the typical "Wassermann" as at present carried out in most laboratories. It may be carried out just as well and with greater economy of material by using one-half the amounts throughout. It is evident that the performance of the reaction calls for experience of serum technique, and knowledge of such reactions, so that fortuitous irregularities may be intelligently controlled. It is our opinion that the performance of routine Wassermann tests by workers without a thorough knowledge of the fundamental facts of

serum phenomena is worse than useless in that insufficient attention to special conditions and to details may easily result in a positive reaction when syphilis is not present, and vice versa.

Recently Archibald McNeil and others have exposed the mixtures of complement, antigen, and patients' serum at refrigerator temperature for a number of hours instead of in the water bath or thermostat at 37.5° C., before adding the sensitized cells. It is a curious fact, which has not yet been satisfactorily explained, that such a procedure increases the delicacy of the reaction. It may be that, when the tubes containing the antigen, patient's serum, and alexin are left at incubator temperature, partial alexin fixation only can take place during the brief period of 30 minutes to one hour, which is usually employed. More prolonged exposure at this temperature would not be advisable on account of deterioration of the alexin. On the other hand, at ordinary ice-box temperatures of about 8° to 10° C., the exposure can be continued for as long as 10 hours without extensive complement deterioration, and meanwhile more complete fixation can occur. This, however, is a surmise. The actual conditions are not clear. As a matter of fact in our laboratory Dr. Ottenberg, in 120 cases so far done in parallel series, one being exposed for fixation for 30 minutes at 37.5° C., the other at 8° to 10° C. for three hours, found discrepancies between the two methods in 15 cases. In all of these, positive reactions were obtained by the ice-box method, whereas by the water bath method the results were negative. Of these cases 7 were clearly unquestionable syphilitics, two were treated syphilis, and four were probably syphilitic.

Many modifications of the Wassermann test have been suggested. Probably the most important is that of Noguchi. The chief justification for this modification is the fact that many normal human sera contain hemolysins for sheep corpuscles. For this reason many workers carry out the ordinary Wassermann technique without adding antisheep sensitizer or amboceptor until they have first observed whether or not the tested serum (in the "back row," without antigen) will not hemolyze the corpuscles without such an addition, adding the sensitizer only when this does not take place. This is advisable since the presence of any considerable amount of normal antisheep sensitizer in the human serum which is being examined (if added to the amount used in the ordinary reaction, 2 units), may so increase the total quantity that hemolysis will result even after most of the alexin has been fixed. Noguchi excludes this uncertainty by avoiding the use of the "sheep cell-antisheep sensitizer" system entirely, substituting a hemolytic complex consisting of human cells and anti-human sensitizer, produced by injecting washed human corpuscles into rabbits.

His technique may be best illustrated in the following tabulation:

Reagents

1. Sensitizer prepared by injecting washed human blood corpuscles into rabbits.
2. 1 per cent. emulsion of washed human blood cells.
3. Alexin—fresh guinea pig serum diluted with one and one-half volumes of salt solution, 40 per cent.

The reaction is performed in the following way:

Noguchi's Method of Complement Fixation for the Serum Diagnosis of Syphilis

| | Set for diagnosis Test with the serum in question | Positive control set Test with a positive syphi- litic serum | Negative control set Test with a normal serum | |
|-----------|---|---|---|---|
| Rear row | a. Unknown serum, 1 drop* b. Complement, 2 units ● c. Corpuscle susp., 1 c. c. | a. 'Positive syph. serum, 1 drop* b. Complement, 2 units ● c. Corpuscle susp., 1 c. c. | a. "Normal serum, 1 drop* b. Complement, 2 units ● c. Corpuscle susp., 1 c. c. | |
| Front row | a. Unknown serum, 1 drop* b. Complement, 2 units ● c. Corpuscle susp., 1 c. c. + Antigen | a. 'Positive syph. serum, 1 drop* b. Complement, 2 units ● c. Corpuscle susp., 1 c. c. + Antigen | a. "Normal serum, 1 drop* b. Complement, 2 units ● c. Corpuscle susp., 1 c. c. + Antigen | Incubation at 37° C. for 1 hour. Addition of antihuman amboceptor, 2 units to all tubes. Incubation at 37° C. for 2 hours longer, then at room temperature. |

* When working with inactivated serum 4 drops (0.08 c. c.) should be employed. With cerebrospinal fluid, 0.2 c. c. (not inactivated) is used.

(Taken from Noguchi's "Serum Diagnosis of Syphilis," Lippincott, 1910, p. 57.)

Bauer³⁰ has introduced a modification in which he utilizes the presence of normal sheep sensitizer in many human sera. He performs his tests without the addition of antisheep sensitizer at first, adding this only to those tubes in which controls have shown that no normal sensitizer is present. Stern,³¹ on the other hand, utilizes the alexin normally present in human serum. The syphilitic serum to be tested is, therefore, not inactivated, and the sheep cells are more heavily sensitized (9 to 12 units). It seems to us that this method is objectionable chiefly because of the anticomplementary action which develops in most normal human sera if kept for a short time, and which can be removed only by inactivation.

Other modifications of the Wassermann reaction are those of

³⁰ Bauer. *Semaine Medicale*, 28, 1908.

³¹ Stern. *Zeitschr. f. Imm.*, Vol. 1, 1909.

Jacobaeus³² and of Wechsleman.³³ It seems, however, that, as the reaction is gaining in importance in clinical diagnosis, most laboratories are adhering to the original system used by Wassermann and his associates, except for the substitution of the non-specific lipoidal antigens for the originally employed organ extracts.

The value of the Wassermann test in the diagnosis of the various stages of syphilis is a problem which can be approached only by careful statistical analysis of the results obtained. This has been done by various investigators, and some of the results have been tabulated in the books of Noguchi, of Boas, and of McIntosh and Fildes. The figures we cite are those largely taken from Boas, as summarized in F. C. Wood's "Chemical and Microscopical Diagnosis" (D. Appleton & Co., 1911), pp. 706 et seq.

Primary syphilis, 974 cases, 56.5 per cent. positive.

The reaction may appear before the primary sore, but this is very rare. Usually it is positive in from 5 to 6 weeks after infection.

Secondary syphilis, 2,762 cases, 88 per cent. positive. In untreated cases they are stated to be 100 per cent. positive.

Tertiary syphilis, 830 cases, 80 per cent. positive.

Tubes, 360 cases, 70 per cent. positive.

Dementia paralytica, 95 to 100 per cent. positive.

The tabulation on the following page, taken directly from Boas, will give a comprehensive summary of this phase of the problem.

Since the reaction is not a specific antigen-antibody union but depends on some substance liberated or produced by reason of the syphilitic injection, it is not out of question that other infections may give rise to a "positive Wassermann." And this, indeed, is the case. It was claimed for a time that a positive reaction may be obtained in tuberculosis, but this has been refuted by subsequent experience, and the earlier positive results probably depended upon faulty technique. There can be little doubt, however, that occasional positive reactions are obtained in cases of leprosy, scarlet fever, malaria, and trypanosoma infections.

The spinal fluid may be used instead of the blood serum in cases of syphilis of the central nervous system, but even here, as Citron³⁴ has shown, the results with blood serum are more frequently positive than those done with the spinal fluid itself. In isolated cases positive reactions have been obtained with ascitic fluids, pleural and pericardial exudates. Bab³⁵ reports a case of positive reaction in

³² Jacobaeus. *Zeitschr. f. Imm.*, Vol. 8, 1911.

³³ Wechsleman. *Zeitschr. f. Imm.*, Vol. 3, 1909.

³⁴ Citron. *Deut. med. Woch.*, 1907, No. 29, p. 1165.

³⁵ Bab. *Münch. med. Woch.*, Vol. 46, 1907.

Table Compiled by Boas, *loc. cit.*, p. 138.

| Stage of disease | Number of cases | Positive reaction | Negative reaction |
|--|-----------------|-------------------|-------------------|
| <i>Control cases</i> (not syphilitic)..... | 1,064 | 1 (scarlatina) | 1,063 |
| <i>Induration</i> | 76 | 56 | 20 |
| <i>Secondary</i> | | | |
| Early untreated..... | 269 | 269 | 0 |
| Recurrent after treatment..... | 199 | 187 | 12 |
| <i>Tertiary</i> | | | |
| No treatment of early tertiary manifestations..... | 63 | 63 | 0 |
| Treatment..... | 20 | 16 | 4 |
| <i>Latent syphilis</i> | | | |
| Within 3 yrs. after infection..... | 243 | 89 | 154 |
| After 3 yrs..... | 111 | 44 | 87 |
| <i>Tabes</i> | | | |
| Untreated..... | 17 | 17 | 0 |
| Treated..... | 26 | 11 | 15 |
| <i>Dementia paralytica</i> | | | |
| Serum..... | 139 | 139 | 0 |
| Spinal fluid..... | 67 | 61 | 6 |
| <i>Congenital</i> | | | |
| With symptoms..... | 54 | 54 | 0 |
| Without symptoms..... | 10 | 7 | 3 |

the milk of a syphilitic mother. Serum obtained at autopsy is not suitable for the reaction, since this, for unknown reasons, may often give a positive reaction in non-syphilitic cases.

COMPLEMENT OR ALEXIN FIXATION AS A METHOD OF DETERMINING THE NATURE OF UNKNOWN PROTEIN

FORENSIC ALEXIN FIXATION TESTS

Our preliminary discussions of the principles underlying alexin or complement fixation have revealed that alexin is bound not only by sensitized cells but also by the specific precipitates formed when an unformed protein antigen is mixed with its specific antiserum. This discovery, made by Gengou, was attributed by him, it will be remembered, to the presence of "albuminolysins," or protein sensitizers, antibodies which have been by many observers regarded as separate from the precipitins, but which we believe, for stated reasons (see p. 193), to be very probably identical with the precipitating antibodies or precipitins. However this may be, when a dissolved antigen is mixed with its antiserum alexin fixation is exerted by the

complex, and this, even when the reacting quantities, antigen and antibody, are so small that visible precipitation will not take place. For this reason, it is plain, it should be possible by means of complement fixation to detect amounts of a foreign protein too small to be demonstrable by direct precipitation with an antiserum.

The method has, therefore, been suggested chiefly by Neisser and Sachs³⁶ for the forensic determination of unknown proteins, as an adjuvant to, and improvement upon, the forensic precipitin test. Our discussion of the principles involved in the introductory paragraphs of this chapter will render unnecessary an extensive discussion of the reasoning upon which this reaction is based. It is well to remind the reader, however, of the facts which we have discussed regarding the quantitative proportions which govern the occurrence of precipitation when an antigen, say human serum, is mixed with its antibody, in this case antihuman rabbit serum. The actual precipitation may be absent either when an excess of the antigen is used or when the antigen is present in too small a quantity. Thus a given quantity of the antiserum may precipitate strongly dilutions of the antigen ranging from 1-50 to 1-10,000. No precipitation or, at least, a very slight one only may occur when concentrations stronger than 1-50 are used and when the dilution is greater than 1-10,000. Nevertheless, in both cases, alexin fixation may be exerted by the complex although no precipitation takes place. As Gay³⁷ has shown, complement fixation may be exerted even when a formed precipitate has been redispersed by the subsequent addition of more antigen. The importance of the forensic reaction of Neisser and Sachs, however, lies chiefly in its application to the detection of quantities of unknown protein too small to be detected by precipitin reactions.

The tests are carried out by mixing a dilution of unknown protein with given quantities of antiserum, adding small quantities of alexin (quantities determined best by previous alexin titration as indicated in our section on the Wassermann reaction); these reagents are left together for a given time at 37.5° C., and then sensitized cells are added to determine whether or not the alexin has been bound.

The table on the following page, taken directly from the article of Neisser and Sachs, loc. cit., will not only illustrate the method of carrying out the reactions but will also give an indication of their extreme delicacy.

It will be seen that 0.00001 c. c. of the normal human serum still gave almost complete complement fixation of 0.05 c. c. of complement in the presence of 0.1 c. c. of the antihuman serum. The table also shows that this reaction follows a general law of relative specificity so often noted in other reactions, namely that, of all the animals tested, the serum of monkeys alone gave reactions with the

³⁶ Neisser and Sachs. *Berl. kl. Woch.*, Vol. 42, No. 44, 1905, p. 1388.

³⁷ Gay. Univ. of Cal., "Publications in Pathology," 1912.

Table Taken from Neisser and Sachs, loc. cit., p. 1388

0.1 human antiserum + 0.05 complement and variable amounts of different normal sera (brought to 1 c. c. volume with salt solution); the mixtures kept 1 hour at room temperature. Then added 1 c. c. 5 per cent. washed beef blood + 0.0015 c. c. amboceptor and left 1-2 hours at 37° C.

The results are as follows:

| Amounts of normal serum | Hemolysis on addition of serum of: | | | | | | | |
|----------------------------------|------------------------------------|----------|----------|----------|----------|----------|----------|----------|
| | Man | Monkey | Rat | Pig | Goat | Rabbit | Ox | Horse |
| 0.01 | 0 | 0 | complete | complete | complete | complete | complete | complete |
| 0.001 | 0 | 0 | | | | | | |
| 0.0001 | 0 | moderate | | | | | | |
| 0.00001 | slight | complete | | | | | | |
| 0.000001 | complete | complete | | | | | | |
| 0 | complete | complete | | | | | | |

human antiserum; and this in quantities as small as 0.001 cubic centimeter.

The forensic complement fixation reaction of Neisser and Sachs is both theoretically and practically valid. Its extensive use in many investigations for theoretical purposes has well established its reliability. However, it is more complicated and requires much more experimental training and care than does the simpler precipitin test, and it will rarely occur that an unknown protein is available in quantities too small to permit of successful precipitation.

THE USE OF COMPLEMENT FIXATION TESTS IN THE DIAGNOSIS OF MALIGNANT NEOPLASMS

A great many attempts have been made to establish a method of complement fixation by which a diagnosis of malignant tumors could be made. It had been hoped that the substance of malignant tumors might contain a form of protein or protein lipid combination which might represent substances specific for such tumors, and might therefore functionate as a specific antigen. On this basis it might be possible that the serum of tumor patients would contain a specific antibody which could react with a specific antigen in tumor extracts, with the resulting formation of an alexin-fixing complex.

No experimental facts have so far justified our assumption of the presence of either specific antigen in tumor extracts, or that of a specific antibody in the serum of such patients. However, we have seen that the Wassermann reaction is a perfectly useful clinically diagnostic method, in spite of the fact that the antigen need not be specific, and the purely empirical basis on which the syphilis reac-

tion is at present based has justified extensive attempts to establish an analogous empirical method for tumor diagnosis.

The literature on this question is confusing. A number of observers using antigens variously prepared from tumor substances have reported favorable results. Simon and Thomas³⁸ report many positive reactions, as do Sanpietro and Tesa,³⁹ and a number of others. Clowes⁴⁰ has carried out a reaction on sarcoma rats and obtained positive reactions in animals in which the tumors were small, negative ones when the tumor had grown to a large size. Ranzi, on the other hand, obtained negative results throughout. Ranzi⁴¹ found that normal serum would often give complement fixation with carcinoma extracts, also that many tumor extracts and sera of tumor patients inhibited complement by themselves. The reactions were so irregular that he assumed them to be without value. Recently the subject has been very thoroughly investigated by v. Dungern.⁴²

Von Dungern claims to have finally evolved a method by which the diagnosis of malignant disease can be made with reasonable accuracy. Like the Wassermann reaction his method is purely empirical. He admits that probably it is not a specific antibody determination and depends rather upon the presence of pathological products of metabolism in the sera of tumor patients. The reliability of his method depends upon the observation of a number of details which he has determined empirically.

He obtains his antigen in a purely non-specific manner, using, as just stated, for this reaction acetone extracts of human blood cells. We take the description of the reaction entirely from his own article in "Weichhardt's Jahresbericht." The antigen is prepared in the following way: Blood is taken from a vein, preferably from a paralytic patient, since v. Dungern claims that individual specimens of blood vary, and he has had the best results with that of paralytic cases. Clotting is prevented by sodium oxalate and the blood cells are thoroughly washed in the centrifuge. To the sediment are added 19 volumes of pure acetone (Merck). This is allowed to stand three days at room temperature and is occasionally shaken during this time. It is then filtered, the acetone evaporated in the incubator at 37° C., and the residue taken up in 96 per cent. alcohol. This alcoholic extract is diluted before use with four parts of salt solution. Of this final preparation 0.8 c. c. is used in the individual test.

Particular precautions must also be taken in the handling of the serum of the patient. In his earliest tests v. Dungern determined

³⁸ Simon and Thomas. *Journ. Exp. Med.*, Vol. 10, 1908.

³⁹ Sanpietro and Tesa. Cited from v. Dungern in "Weichhardt's Jahresbericht," etc., Vol. 8, 1912, p. 163.

⁴⁰ Clowes. *Journ. A. M. A.*, 1909, Vol. 52.

⁴¹ Ranzi. *Wien. kl. Woch.*, 1906, p. 1552.

⁴² V. Dungern. *Münch. med. Woch.*, Nos. 2, 20, 52, 1912; *Berl. kl. Woch.*, 1913, "Weichhardt's Jahresbericht," Vol. 8, 1912, p. 163.

that the inactivation of the tumor sera greatly diminishes their specific fixation properties, and for this reason he at first advised that the serum be used unheated. He has found recently that the best results are obtained when the serum is heated to 54°C. , together with a little sodium hydrate solution. He handles the blood in the following way: After being taken from the patient it is allowed to stand 1 to 2 days in the refrigerator; just before use he adds two parts of an $\frac{N}{50}$ NaOH solution with one part of serum and heats it for half an hour at 54°C. As it is important that the sodium hydrate should contain no sodium carbonate, he advises the use of the Kahlbaum preparation. In setting up the test he uses graded quantities of the mixture corresponding to 0.2, 0.1, 0.05, and 0.025 c. c. of the original serum. To each of these quantities he adds the stated quantity, 0.8 c. c. antigen preparation described above, and the 0.05 guinea pig complement. Controls must be set up with the antigen alone and with the patient's serum alone to prevent error from independent fixation by these substances. These reactions are allowed to stand three hours at room temperature, and then one cubic centimeter of a 5 per cent. solution of beef blood sensitized with two units of hemolytic serum is added (as in the Wassermann reaction). It is important to use a strongly sensitizing serum, so that not too much of the hemolytic rabbit serum must be added to the tubes. Experiments done in this way with normal sera usually result in complete hemolysis within one hour, although in certain other diseases, i. e., tuberculosis and syphilis, slight inhibition may result. However, fixation with the patient's serum in quantities of 0.1 c. c. or less is, according to von Dungern, fairly specific for malignant tumors, since normal sera treated in the way described usually do not cause fixation in quantities of less than 0.2 c. c. and, in syphilis and tuberculosis, if fixation is at all present, it is usually not evident in quantities less than 0.1 c. c.

With a reaction so carried out von Dungern has examined 244 cases. The following tabulation states his results:

| Malignant tumor of | No. of cases | Reaction positive* |
|--------------------|--------------|--------------------|
| Pharynx..... | 3 | 3 |
| Esophagus..... | 6 | 6 |
| Stomach..... | 15 | 11 |
| Rectum..... | 14 | 10 |
| Larynx..... | 2 | 2 |
| Tongue..... | 5 | 5 |
| Bladder..... | 1 | 1 |
| Breast..... | 22 | 22 |
| Uterus..... | 10 | 10 |
| Skin..... | 8 | 7 |
| Ethmoid bone..... | 1 | 1 |
| Upper maxilla..... | 1 | 1 |

* Taken from von Dungern, "Weichhardt's Jahresbericht," Vol. 8, 1912, p. 174.

We report von Dungern's results exactly as he states them in his last summary, since his well-known experimental ability necessitates serious consideration of all of his work. We may say, however, that a survey of the entire literature of complement fixation in the diagnosis of malignant tumors does not yet justify our acceptance of this method as of anything like the established value which the similar method has attained in syphilis.

COMPLEMENT FIXATION IN GLANDERS

The diagnosis of glanders by the mallein test and by agglutination has been recently reënforced by the method of complement fixation. In carrying out these tests the method of preparation of the antigen is of the greatest importance. The directions which we give are those employed in the Diagnostic Laboratory of the New York Department of Health, under the immediate supervision of Dr. McNeil and Miss Olmstead, from whom we have our information.

The particular strain of glanders bacilli employed seems to be of little importance. The organisms are grown on 1.6 per cent. acid glycerin-potato-agar. This stock culture is transplanted every other day. From it cultures are planted upon salt-free veal peptone agar. It is of the greatest importance that this medium shall be neutral to phenolphthalein. After twenty-four hours in the incubator the growth is washed off with distilled water, which also should be neutral, and the emulsion heated for from four to six hours at 80° C. in a water bath. It is then filtered through a Buchner filter simply to facilitate subsequent filtration through a Berkefeld "N" or "V" filter. After filtration this antigen is again sterilized for one hour at 80° C. and then on two successive days at 56° C. for one-half hour.

The fixation tests carried out with these antigens have yielded excellent results as reported by Dr. McNeil⁴³ at the New York Serological Society.

It is unnecessary to give further directions as to the technique of this reaction, since it is simply that of complement fixations in general, the chief difficulty being that of antigen preparation.

COMPLEMENT FIXATION IN GONORRHEAL INFECTIONS

There are certain conditions following gonococcus infection of the genito-urinary tract which are not easily distinguished from a number of other tests unless the organisms can be cultivated or a specific serum reaction can be applied. Most important of these are gonorrheal rheumatism, salpingitis, and endocarditis. Complement fixation with the sera of such patients, and an antigen produced from

⁴³ McNeil, Archibald. N. Y. Serological Soc. Meeting, April 4, 1914.

gonococci, has been employed by many observers during recent years, and promises to be of great value.

Here, too, the production of the antigen is the only feature of the reaction which has offered difficulties. Since the researches of Torrey have shown that not all races of gonococcus are antigenically alike, it seems necessary to produce a polyvalent antigen. At the New York Department of Health at present the antigen is prepared by using the ten Torrey strains. Stock cultures are carried on neutral veal agar and, for antigen preparation, cultures are planted upon a salt-free veal agar. Twenty-four-hour growths are washed off in neutral distilled water, are kept in a water bath at 56° C. for two hours, and are then filtered through, first, a Buchner and then a Berkefeld filter. They are then sterilized for one hour. The antigen so prepared is now ready to be titrated and used.⁴⁴

⁴⁴ For information concerning the details in the preparation of this antigen we are indebted to Miss Olmstead of the N. Y. Department of Health.

CHAPTER IX

THE PHENOMENON OF AGGLUTINATION

WHEN bacteria are added to homologous immune serum there occurs a peculiar agglomeration of the individual micro-organisms into small clumps. The phenomenon is so general and so easily observed that it is not surprising that it was noticed and reported by a number of workers during the period of active investigation upon serum reactions which preceded and followed the discovery of the Pfeiffer phenomenon. Thus, in the years from 1891 to 1895, Metchnikoff,¹ Charrin and Roger,² Isaëff and Ivanoff,³ Washburn,⁴ and several other workers made this observation with a variety of bacteria and immune sera. But all of these observers failed to follow up or analyze the process they incidentally noticed in the course of other investigations. A thorough study of the phenomenon was not made until 1896, when Gruber and Durham,⁵ in Vienna, in the course of their studies upon bacteriolytic reactions with colon bacilli and cholera spirilla, again noticed the agglutination of these bacteria in homologous immune sera, recognized the specificity of the reaction, and called attention to its apparent independence of other previously studied serum phenomena.

The process known as agglutination consists in the following train of occurrences. If we add to an even emulsion of bacteria a small amount of homologous immune serum the micro-organisms may be seen to collect rapidly in groups or masses, with a resultant clearing of the fluid in which they have been suspended. The clumps of bacteria gather in flakes which, not unlike flakes of snow, sink to the bottom of the test tube. The speed and completeness with which this phenomenon occurs depend, as we shall see, upon the agglutinating strength and other qualities of the serum which is employed, but the essential process of clumping is alike in all cases.

There are a large number of different methods by which this

¹ Metchnikoff. *Ann. de l'Inst. Past.*, 1892.

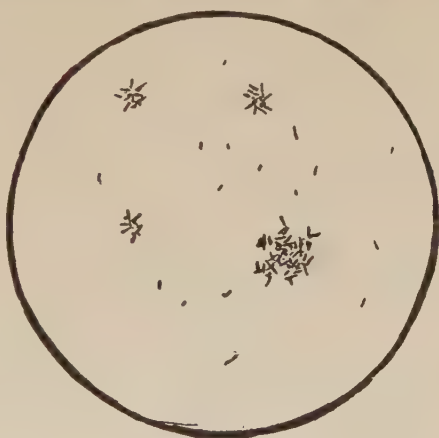
² Charrin and Roger. *C. R. de la Soc. de Biol.*, 1889.

³ Isaëff and Ivanoff. *Zeitschr. f. Hyg.*, Vol. 17, 1894.

⁴ Washburn. *Journ. of Path. and Bact.*, 1896, p. 228.

⁵ Gruber and Durham. *Munch. med. Woch.*, 1896.

occurrence can be observed, each one particularly adapted to some special purpose for which the reaction is carried out. Gruber and Durham, who were investigating the properties of bacteriolysins when they observed agglutination, naturally recognized the specific nature of the reaction and proposed to make use of it for the purpose of bacterial differentiation and species determination. For this purpose, which has become one of the most important of the practical applications of the agglutination reaction, the phenomenon is best observed by the so-called "macroscopic method," in which a series of serum dilutions are mixed, in small test tubes, with equal volumes of emulsions of the bacteria. Thus, if we wish to determine the nature of an unknown bacillus, suspected of belonging to the typhoid bacillus group, by this method, we may determine its agglutination in the serum of an animal immunized with a known strain of typhoid. The tubes are incubated after the mixtures have been made, and the agglutination which has taken place in the various tubes is recognized by a clearing up of the fluid and the flaking of the bacteria after from one to three hours. The test tube method has the advantage of permitting the use of larger quantities of reagents than can be used in the other methods described below, and therefore more exact quantitative measurements can be made.



MICROSCOPIC AGGLUTINATION.

Although this method for the determination of bacteria has found universal application, it is probably most frequently employed at the present time for the rapid identification of colonies of doubtful typhoid or dysentery, incident to the isolation of these organisms for stools by such methods of plating as those of Conradi-Drigalski, of Endo, or of Hiss. The suspicious colonies can thus be fished directly to an agar slant, and the cultures, when developed, emulsified and identified by agglutination. The advantages of such a method for the determination of the biological interrelationship of the organisms of a given group, like, for instance, that of the dysentery bacilli, are obvious.

An ingenious use of this reaction was also made by Shiga when he determined, among various bacteria isolated from the stools of dysentery cases, the particular one which was specifically aggluti-

nated by the patient's serum, thus discovering the dysentery bacillus which bears his name.

Within a few months after the publication of Gruber and Durham's work, Widal and, apparently independently of him, Grünbaum,⁶ by a process of reasoning the converse of that detailed above, applied the reaction to the diagnosis of infectious disease.

It is obvious that a human being or an animal infected with a given variety of bacteria may develop agglutinating properties against these bacteria. It is of great value, therefore, to determine the agglutinating power of the serum of a patient for the bacteria which are known to cause the disease suspected in the particular case in which a diagnosis is desired. This method has become a routine measure in the early diagnosis of typhoid fever under the name of "Widal" or "Gruber-Widal" reaction and, since the quantities of serum which can easily be obtained from a patient are usually small, it is convenient to carry out the reaction by the microscopic method. This consists in mixing serum and bacterial emulsion in hang-drop preparations and observing them with the microscope. An excellent method, also, is the so-called Proescher⁷ method in which serum and bacterial emulsion are mixed in small watch-glasses or salt cellars. Proescher used this method extensively in the study of staphylococcus agglutinations. The mixtures in the salt cellars were set away at 37° C. for two hours, and then observed with a magnification of 60 to 70 diameters.

Close observation of the occurrence under the higher power of a microscope shows that the bacteria, if motile, lose their motility, if non-motile the Brownian motion is arrested. They are then rapidly gathered in small clumps, isolated individuals between these clumps being gradually drawn into them, until finally the fluid between the masses is entirely clear. This complete clearing, of course, happens only when there is not an excess of bacteria, for, like other serum reactions, this phenomenon is a quantitative one in which definite proportions must be maintained.

Clinically the most frequent use of the agglutination reaction is in the diagnosis of typhoid fever. The technique used for this test is, in the large majority of cases, the microscopic hang-drop method. In Germany the Proescher method is sometimes used, and the microscopic method with dead organisms, as first introduced by Ficker, is also not uncommon at the present day.

Since the serum of normal human beings very often contains moderate agglutinating powers for the typhoid bacillus, the diagnostic value of the reaction in this disease depends upon the elimination of this error by sufficient dilution. If dilutions of the serum of from 1-40 to 1-60 are used diagnostic errors on this point are

⁶ Grünbaum. *Lancet*, 1896, Vol. 2.

⁷ Proescher. *Centralbl. f. Bakt.*, Vol. 34, 1903.

avoided, since the normal agglutinating power of human beings is never such that typhoid bacilli will be clumped by it in these dilutions within one hour. Prompt clumping in serum dilutions of 1-20 is fairly reliable, but does not absolutely exclude an unusually high normal agglutinating power. In carrying out tests clinically dilutions of 1-20, 1-40, and 1-80 are usually made and observed for one hour. From such tests diagnosis can be made without danger of error. In rare cases of icterus the agglutinating power for typhoid bacilli may be increased. Just what is the cause of this is not certain; Wood^s reports cases in which agglutination of 1-40 was present with slight jaundice (hepatic cirrhosis). On the other hand he has frequently failed to notice agglutination in other cases of intense jaundice. It is not impossible, as Wood suggests, that the occasional presence of unusual agglutinating power in individuals with jaundice has some relation to the frequent persistence of typhoid bacilli in the gall bladder.

Occasionally it will be noticed that dilutions of the patient's serum of 1-5 to 1-20 fail to agglutinate, while higher dilutions will give positive tests. This is referable to the so-called "pro-agglutinoid zone," the principles underlying which we shall discuss in another place.

The Widal test in typhoid cases rarely appears before the end of the first week, and, in the majority of cases, is present before the end of the second week. It may proceed for months, although Wood states that he has seen it disappear at the end of three to six weeks.

In *paratyphoid fever* the diagnosis can often be made by agglutination, and in *dysentery*, as we have seen, the fact that the patient's blood agglutinated the bacteria was one of the important facts utilized by Shiga in his discovery of the organism which bears his name.

In *pneumonia* agglutination of the pneumococcus, isolated from the patient's sputum by sera prepared by immunization with various types of pneumococci, has become of considerable importance clinically, since Neufeld and Haendel and, in this country, Cole, Dochez, and Gillespie have determined that there are a number of different types of this micro-organism. The use of pneumococcus serum in the disease will be of value only if a serum is used which has been produced with an organism of the same type as the one infecting the patient. Therefore, determinations of the type by highly potent agglutinating sera give a finger-point to the variety of serum to be used. Whatever may be the eventual outcome of the serum treatment in pneumonia, no results whatever can be expected, according to our present knowledge, unless such determinations are made. The technique of agglutinations in pneumococcus work is facilitated by

^s Wood. "Chemical and Microscopical Diagnosis," Appleton & Co., p. 242.

growing mass cultures of organisms, as advised by Hiss, in flasks of glucose broth containing 1 per cent. calcium carbonate.

The same method of growing micro-organisms is useful in the case of streptococcus agglutinations, since the insoluble calcium carbonate, if thoroughly shaken, breaks the chains of streptococci and thereby facilitates judgment as to the reaction.

Agglutination reactions have been of considerable usefulness also in the diagnosis of glanders in horses. The early work on this subject was done chiefly by MacFadyean,⁹ and the reaction has been particularly studied by Wladimiroff.¹⁰ Since the serum of normal horses will often agglutinate glanders bacilli in dilutions of as much as 1-500, Wladimiroff advises making the positive diagnosis on dilutions only higher than 1-1,000, since he states that normal horses may occasionally reach an agglutination titre of 1-1,000. The same writer states, moreover, that glanders bacilli are subject to great variations in agglutinability, and that for this reason the choice of a suitable strain is of great importance.

The motility of bacteria has absolutely no relation to the reaction, and their agglutination is entirely passive.

Some of the earlier investigators of agglutination associated the reaction with alteration in the flagellar mechanism of the micro-organisms. It is now well known, however, that non-motile, as well as motile, bacteria are subject to the phenomenon, and that no visible change in the appearance or arrangement of flagella accompanies the clumping. Although this is the case, observation of the motility of such organisms as the bacillus of typhoid fever, while subjected to the action of agglutinating serum, may be of great value in aiding in the determination of the degree of completeness with which the reaction is taking place.

Agglutination, furthermore, does not lead to the death of the bacteria. Of course, whenever the reaction is carried out in unheated serum the concomitant effects of the bactericidal substances bring about bacterial death. Agglutination does not, however, depend upon the coöperation of alexin, and serum may be inactivated without interference with its power of agglutination. In such heated serum clumping takes place without bactericidal effects, and, more than this, the bacteria may grow, if exposed to proper temperature conditions, when suspended in the serum. In fact, it is of considerable interest to carry out the reaction in this way, for the bacteria growing in agglutinating serum form long convoluted threads and skeins even when in ordinary culture they habitually occur as separate individuals. Thus colon bacilli, typhoid bacilli, pneumococci, cholera spirilla, and other organisms, which ordinarily grow as free single cells, or, at most, in chains of two or three, if kept in the

⁹ Macfadyean. *Journ. of Comparative Path. and Ther.*, Vol. 9, 1896.

¹⁰ Wladimiroff. "Kolle u. Wassermann Handbuch," 2d Ed., Vol. 5.

incubator for ten to twelve hours together with homologous serum, will grow in long, delicate chains, like those of streptococci. This form of reaction has been especially studied by Pfaundler,¹¹ who attributed particularly delicate specificity to it. However, the "Thread Reaction" of Pfaundler, as it is sometimes called, is merely another manifestation of the phenomenon of agglutination and subject to the same laws and limitations of specificity which apply to other methods.

The purely passive rôle played by the bacteria in agglutination is best shown by the fact that dead bacteria, killed in various ways, are specifically clumped just as are the living cultures.¹² On this fact depends the method spoken of as "Ficker's Reaction," in which emulsions of typhoid bacilli, killed by formaldehyd or carbolic acid (distributed commercially), are agglutinated in small test tubes by the serum of typhoid patients. The original method of Ficker is said to be a proprietary secret; however, a number of other methods which attain the same purpose are in use in various places. Volk¹³ describes the method used in Vienna, and states that there carbolic acid is used to kill the cultures. Similar to this is the method described by J. H. Borden,¹⁴ who proceeds as follows:

The bacilli are grown on agar slants in large tubes for 24 hours. They are then washed from the medium with a sterile mixture of salt solution 450 parts, glycerin 50 parts, and 95 per cent. carbolic acid 2.5 parts. After this solution has been kept a week it becomes translucent and by this time the bacilli are dead. The preparation is then ready for use and can be kept a long time in dark bottles in a cool place. Borden very carefully controlled this bacterial emulsion with positive and negative typhoid sera and found it reliable. The great advantage of all these methods, of course, consists in the possibility of furnishing the general practitioner with materials for clinical agglutination tests in which the necessity of preserving and suspending living cultures is eliminated.

The facts which we have just considered tend to show that agglutination is not a vital phenomenon¹⁵ dependent in any way upon the living nature of the bacterial cell, but, like other phenomena of antigen-antibody reactions, a purely chemical or physical process in which the substance of the bacterial cell enters specifically into relation with the agglutinating factor of the serum. In uniformity with other analogous reactions the antigenic substance is here spoken of as "agglutigen," the antibody as "agglutinin."

¹¹ Pfaundler. *Wien. kl. Woch.*, 1898, and *Centralbl. f. Bakt.*, I, Vol. 23, 1898.

¹² Bordet. *Ann. Past.*, Vol. 10, 1896.

¹³ Volk. "Kraus und Levaditi Handbuch," Vol. 2.

¹⁴ Borden. *Medical News*, N. Y., Mar., 1903.

¹⁵ Bordet. *Ann. Past.*, Vol. 10, 1896.

The agglutinin, or agglutinin-inducing substance in the bacteria is apparently an inherent part of the bacterial protein, and agglutinins may be produced in animals by injection not only of living and dead whole bacteria, but by bacterial extracts, prepared in various ways. And, furthermore, just as the agglutinins of serum are absorbed out of a serum by the whole bacteria, they may be neutralized by the dissolved bacterial extracts.

Just what the nature of the agglutinin is has been a matter of prolonged controversy, Pick¹⁶ and others claiming that it is possible to obtain an agglutinin by alcohol precipitation from old bacterial cultures which, upon further purification, can be found to give none of the usual protein reaction (Buiuret, Millon). It is by no means certain, however, that Pick's results are correct. In fact, many objections have been advanced against them, and the acceptance of an antigen of non-protein nature is so opposed to all our knowledge regarding antigens in other cases that Pick's results should not be taken as final until very careful revision of the experimental facts has been carried out. That the agglutinin is, to a certain extent, subject to dialysis has been claimed because of experiments in which specific agglutinins have appeared in the sera of animals into whose peritoneal cavities celloidin sacs, filled with bacteria, have been placed.¹⁷

There has been a great deal of discussion regarding the possible localization of the agglutinin of bacteria in the ectoplasmic layers of the cells, and especially in the flagellar substance. We have seen that, as a matter of fact, nonmotile bacteria are subject to the phenomena of agglutination just as are the motile forms, but numerous attempts were made during the earlier stages of our knowledge of this reaction to demonstrate that changes in ectoplasm and flagella accompanied the actual agglutination. Gruber¹⁸ himself held the opinion for a time that the clumping was due to an ectoplasmic swelling which rendered the bacteria sticky, causing them to hold together after chance approximation. He soon gave up this idea himself, but a similar theory was for some time upheld by Nicolle¹⁹ and others.

Malvoz²⁰ in 1897 devised an ingenious method by which he believed that he could show that the agglutination of bacteria depended upon their ectoplasmic substances. He passed the typhoid emulsion through Chamberland filters and, when the bacilli had been caught

¹⁶ Pick. "Hofmeister's Beiträge," 1901, 1902.

¹⁷ This would be in keeping with Pick's work just referred to, and should be subjected to the same criticism before final acceptance. For a more detailed discussion of these conditions the reader is referred to the article by Paltauf, "Kolle u. Wassermann Handbuch," Vol. 4, part 1.

¹⁸ Gruber. *Münch. med. Woch.*, 1896.

¹⁹ Nicolle. *Ann. de l'Inst. Past.*, 1898.

²⁰ Malvoz. *Ann. de l'Inst. Past.*, Vol. 11, 1897.

upon the filters, he subjected them to prolonged washing. The bacilli, now removed from the filter by passing fluid through in the opposite direction, were no longer motile or agglutinable either by formalin, safranin, or other chemical agents, nor by agglutinating sera. Dineur,²¹ repeating the experiments of Malvoz, came to the same conclusions. He decided that in agglutination the bacteria became entangled with each other by means of the flagella. Harrison,²² in later studies working under Tavel, attempted to dissolve out the ectoplasmic layers of bacteria with pyocyanase, and from his experiments also came to the conclusion that the agglutininogen was contained in the external layers. Similar results were obtained by De Rossi.²³

Further studies on the same problem are those of Smith and Reagh.²⁴ These investigators worked with two strains of bacilli, both of which they regarded as belonging to the hog-cholera group, though the one was motile and the other nonmotile. When rabbits were immunized with the nonmotile bacillus an agglutinin was obtained which acted upon this bacillus differently and less powerfully than did the agglutinin produced with the motile one. Contact with the nonmotile bacillus did not deprive the serum produced with the flagellated organism of the agglutinins for the latter. They conclude that two agglutinins were involved—one incited by the ectoplasm and flagellar substance, the other by the bacterial cell body proper. Rehns as well as Paltauf have criticized these results by questioning the species identity of the two bacilli employed in the experiments, referring the phenomenon to the occurrence of group agglutination.

As a matter of fact our present knowledge of agglutination no longer justifies the association of agglutination with flagella. Non-motile as well as motile bacteria are readily agglutinated, and we have much evidence which will be discussed presently which shows that the agglutination reaction is governed by many of the laws which obtain in colloidal flocculations. This, however, does not exclude the possibility that it is the ectoplasmic zone chiefly which takes part in the reaction. Furthermore, loss of motility, which always accompanies agglutination when a motile organism is under observation, is an extremely valuable aid in guiding us in our judgment of incomplete reactions.

That changes may be brought about in bacterial agglutininogen by various methods of treatment has been shown by a number of work-

²¹ Dineur. *Bull. de l'Acad. de Méd. de Belge*, 1898, cited from Smith and Reagh.

²² Harrison. *Centralbl. f. Bakt.*, Vol. 30, I, Orig. 1901.

²³ De Rossi. *Centralbl. f. Bakt.*, I, Vols. 36 and 40.

²⁴ Smith and Reagh. *Journ. of Med. Res.*, Vol. 10, 1903.

ers, although the fundamental principles underlying such changes are not at all clear.

Joos²⁵ was the first to study agglutination with particular reference to the effects upon the reaction of heating both the antigen and the antibody. On the basis of extensive and complicated experiments upon the agglutinin produced in horses by immunization with heated and unheated typhoid bacilli, he drew the conclusion that agglutinin (agglutinin-inducing substance) in bacteria was not a single element but consisted of at least two definite parts of which he speaks as α and β -agglutinin. α -agglutinin is a constituent of the living bacteria, and is easily destroyed at 60° to 62° C. The β -agglutinin is also present in normal bacilli, but is more heat-resistant and will withstand 60° to 62° C. for several hours. The injection of living unheated bacilli then induces the formation of both α and β -agglutinin, which have respectively a particular affinity for α and β -agglutinogens. The injection of heated bacilli, on the other hand, induces the formation only of β -agglutinin and not a trace of α -agglutinin. The α -agglutinin is considerably heat-resistant, resisting 60° to 62° C., whereas the β -agglutinin loses its agglutinating property when heated to 60° C. The α -agglutinin is entirely incapable of uniting with β -agglutinin. However, β -agglutinin can combine or react with both the α and β constituents of the bacilli. For this reason Paltauf has spoken of agglutinin produced with the heated bacteria as "umfänglicher." This is a point of great interest, and if Joos is right is, of course, of considerable practical importance.

However one may look upon these experiments, as well as the similar ones of other workers, it seems established that heating bacteria leaves them still capable of inciting agglutinins powerfully and rapidly, perhaps of an "umfänglicher" nature than those produced with the native cells.

Heating bacteria may also alter their agglutinability. Thus, according to Eisenberg and Volk,²⁶ heated above 65° C. the bacteria no longer agglutinate in the presence of specific immune serum, but still absorb agglutinin. Eisenberg and Volk, therefore, distinguish between a heat-sensitive constituent of the antigen, which is particularly associated with the clumping, whereas the thermostable substance represents the haptophore or combining portion. It seems simpler, in this case also, to assume a change in the colloidal stability of the bacteria by heating than to seek it in a differentiation into combining and agglutinable parts of the same antigen.

The points raised by Joos' work have been followed up particu-

²⁵ Joos. *Centralbl. f. Bakt.*, Vol. 33, 1903.

²⁶ Eisenberg and Volk. *Zeitschr. f. Hyg.*, Vol. 40, 1902.

larly by Kraus and Joachim²⁷ and by Scheller.²⁸ Scheller summarizes the results of his work as follows: First, in agreement with Joos he found that immune sera obtained by injection of bacteria modified by heat vary considerably from each other. Secondly, immunization with living typhoid bacilli produces sera which agglutinate living bacilli very highly and less highly bacilli heated to 60° C. The titre of agglutinating serum is altered very little toward living bacilli after heating to 60° to 62° C., but is sometimes diminished toward bacteria that have been heated. Bacilli that have been heated to 100° C. but slightly agglutinate unheated serum. Sera produced by the injection of typhoid bacilli heated to 60° to 62° C. agglutinate with both living and heated bacilli. Very important furthermore in Scheller's work are the determinations that typhoid bacilli which have been heated absorb agglutinins out of the sera more actively than do the unheated bacteria, and that the highest agglutinin titres can be obtained by agglutination with bacilli that have been heated to 60° C. The analogy of Scheller's results with similar work done in connection with the precipitin reaction is striking and will be referred to in another place.

Alterations in the agglutinability of bacteria may also occur spontaneously, without previous heating, as in the preceding experiments. It has been frequently noticed that typhoid bacilli, recently cultivated out of the human body, will not agglutinate in sera which have high agglutinating power for strains kept for some time on laboratory media. Much investigation has been focused upon the determination of the cause for this, and although many explanations have been suggested no satisfactory solution has been found. Most workers who have attempted to attack this problem have based their reasoning upon the receptor conception of Ehrlich and have assumed that such inagglutinable bacteria are characterized by a diminished equipment in "receptors." Such strains have been especially well studied in the case of typhoid bacilli and cholera spirilla. Inagglutinable typhoid bacilli have been cultivated by many investigators from the spleen, gall bladder, and urine of typhoid patients, and many of these, when studied for prolonged periods, have been found to regain normal agglutinability after several generations of cultivation upon artificial media. Apparently some alteration of the bacilli had taken place in the presence of the body fluids (immune serum) which affected their sensitiveness to the agglutinins, i. e., their ability to unite with or absorb this antibody. The phenomenon involves an important principle, emphasized some years ago by Professor Welch, namely, that the bacteria may acquire a quasi-immunity against the attacking forces of the body, a property which may be responsible for the increase of virulence noted when some

²⁷ Kraus and Joachim. *Centralbl. f. Bakt.*, I, Vol. 36, 1904.

²⁸ Scheller. *Centralbl. f. Bakt.*, Vol. 36, 1904, and Vol. 38, 1905.

bacteria are repeatedly passed through the bodies of animals, and, indeed, alterations of virulence signify biologically a process of adaptation on the part of the bacteria just as increased immunity indicates a similar process on the part of the invaded subject.

This lessened susceptibility to antibodies is noticeable not only in strains cultivated from the body in disease, but can be produced artificially by cultivating the bacteria in inactivated homologous immune serum. This has been accomplished by Walker²⁹ especially, and by Müller,³⁰ with both typhoid bacilli and cholera spirilla cultivated upon broth mixed with serum. Such strains not only increase in virulence but lose in both agglutinability and susceptibility to bactericidal effects. Sacquépée³¹ obtained similar results by keeping the organisms in collodium sacs in the peritoneal cavity, and Bail³² found similar inagglutinability of typhoid bacilli taken from the peritoneal exudates of guinea pigs dead of infection.

Zinsser and Dwyer³³ have noticed similar inagglutinability in typhoid bacilli recovered from the peritoneal cavities of guinea pigs injected with anaphylatoxin and bacteria. The anaphylatoxin in these cases possessed distinct aggressive action, and the conditions here were probably very similar to those observed by Bail.

There are various possible explanations, the most prevalent ones all representing variations of the opinion that such inagglutinable strains possess an inadequate receptor apparatus. Cole³⁴ advances this because he found these cultures possessed less power to absorb agglutinin than others, and, injected into animals, produced sera which were not highly agglutinating for the injected strain. Some of Cole's experiments show clearly the variable agglutinability displayed by different strains of the same species. Thus the agglutination in the same serum

| | |
|---------------|---------|
| Of strain E = | 1:8,000 |
| Of strain H = | 1:7,000 |
| Of strain I = | 1:4,500 |
| Of strain W = | 1:4,500 |
| Of strain C = | 1:4,000 |

The difference here between E and C actually amounted to a relation of 1 to 2. A rabbit immunized with strain I furnished a serum which agglutinated strain E more powerfully than I itself.

Müller's experiments have the same general significance. It has also been suggested that the inagglutinable bacteria, especially those from the peritoneal exudate, which Bail found were neither agglu-

²⁹ Walker. *Journ. of Path. and Bact.*, Vol. 8, 1902.

³⁰ Müller. *Münch. med. Woch.*, 1903.

³¹ Sacquépée. *Ann. Past.*, Vol. 4, 1901.

³² Bail. *Archiv f. Hyg.*, Vol. 42.

³³ Zinsser and Dwyer. *Proc. Soc. for Exp. Biol. and Med.*, Feb., 1914.

³⁴ Cole. *Zeitschr. f. Hyg.*, Vol. 46, 1904.

tinable nor absorbed agglutinin, may have taken up altered agglutinin or agglutinoid. We will have occasion to recur to this problem in connection with our discussions of the capsulated bacteria and of virulence. The explanations given above do not seem on the whole satisfactory, and the problem is an exceedingly complex one. It has been found indeed that the acquired resistance of bacteria against agglutinins is not at all unique, and that acquired resistance against serum lysins may be observed.³⁵ The extensive investigations of Bail, Walker,³⁶ and others, on the nature of changes in virulence in many invasive bacteria, and the knowledge more recently gained on the resistance to phagocytosis of virulent strains of streptococci and pneumococci are facts closely related in underlying principle to the inagglutinability of typhoid strains cultivated in immune sera.

That no two strains of bacteria of the same species are exactly similar in their agglutinability in the same serum, moreover, is an observation which is made by every one who is in a position to carry out routine Widal tests in hospital practice. The spontaneous agglutination which occasionally occurs in the broth cultures of typhoid bacilli used for this test in many laboratories³⁷ may often be referred, at least in the cases which have come to the writer's notice, to an excessive acidity of the broth, a phenomenon which will be discussed in a subsequent paragraph. As far as the phenomenon of variable agglutinability inherent in different strains is concerned, however, it is of great practical importance in carrying out routine Widal tests to bear this in mind and to control the strain of typhoid bacilli employed in the reactions from this point of view. A strain also which has been in use for a long time should be titrated with agglutinating animal sera from time to time to determine whether or not alterations in agglutinability have occurred.

That the reaction of bacterial agglutination was specific was noted, we have seen, by Gruber and Durham from the very beginning. The closer study of the reaction in its application to bacterial identification has led to interesting data which have revealed certain definite limitations of this specificity. It has been found, for instance, that, while immunization with any given species of bacteria gives rise to a very marked increase of agglutinins for this species, there are formed at the same time, though to a lesser degree, agglutinins for bacteria of other species. This has been referred to as "group reaction," and the agglutinins appearing in such sera are spoken of by German observers as "*Haupt Agglutinine*" and "*Neben*" or "*Mit Agglutinine*." In English texts they are usually referred to as "*chief*" or "*major*" agglutinins and "*para*" or "*minor*" agglutinins. Although, as a general rule, such group-agglutinin formation

³⁵ Eisenberg. *Centralbl. f. Bakt.*, Vol. 34, p. 739, 1903.

³⁶ Walker. *Centralbl. f. Bakt.*, Vol. 33, 1903.

³⁷ See section on Aggressins.

is evident most markedly in the cases of biologically related micro-organisms like the typhoid, paratyphoid, and colon bacilli, this is not necessarily the case, and in some instances the immunization of an animal with a given bacterium may produce minor agglutinins for other bacteria that have no morphologically or culturally demonstrable biological relation to that which reacts with the major agglutinin. We may obtain the most graphic survey of these conditions by examining one of a number of experimental protocols in which such major and minor agglutinin formation is illustrated. Thus in the work of Hiss³⁸ on the dysentery bacilli the following relations were observed:

A serum produced in rabbits by immunization with the Shiga bacillus agglutinated the Shiga bacillus in dilutions of 1 to 20,000, the "Baltimore," "Harris," "Gray," and "Wollstein" bacillus 1 to 1,200, the Y bacillus and others 1 to 200.

An Anti-Y bacillus serum, which agglutinated this bacillus 1 to 6,400, agglutinated the Baltimore bacillus 1 to 1,600, and the Shiga bacillus 1 to 100.

Anti-"Baltimore" bacillus serum agglutinated this bacillus 1 to 3,200, and the Y bacillus 1 to 400, and the Shiga bacillus 1 to 100.

In this series there is fair correspondence between cultural biological relations and agglutination, and from many such investigations it would seem that "group" agglutination might be taken to represent a method of determining biological classifications similar to the zoölogical relations revealed by the precipitin reaction. While, in a general way, this is undoubtedly true, nevertheless great caution must be exercised in relying upon such evidence for classification, since notable exceptions have been observed. Park,³⁹ for instance, cites a case in which a horse, immunized with a paradysentery bacillus, agglutinated a colon strain in dilutions up to 1 to 10,000. Similarly Durham⁴⁰ found that two members of the colon group—one saccharose fermenting—reacted almost identically with the same agglutinating serum, while the agglutinations of two culturally identical bacilli of the hog cholera group were entirely at variance.

The cause for the phenomenon of group agglutination must unquestionably be sought in the nature of the bacterial agglutinogens, and it is but reasonable to assume that living cells so little differentiated biologically and morphologically should have much in common chemically as well. The bacterial cell, moreover, may contain several antigenic complexes and, beside its specifically peculiar constituents, therefore, we may suppose that every bacterium contains additional antigenic substances which it has in common with other

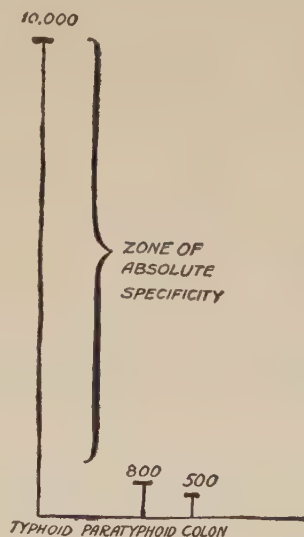
³⁸ Hiss. *Journ. of Med. Res.*, 13, N. S., Vol. 8, 1904.

³⁹ Park. "Pathogenic Micro-organisms," 1910, p. 166.

⁴⁰ Durham. *Journ. of Med. Res.*, Vol. 5.

species. It is the specific antigen in response to which the "chief" agglutinin is formed, while the others, present in smaller quantity, lead to the formation of the minor or paraagglutinins with an intensity proportionate to the amounts present in the bacterial cell. Thus, as Durham expresses it, if we assume one micro-organism to contain antigenic substances a, b, c, and d, and another d, e, f, and g, the antibodies produced by injections of the former would react with the common element d in the latter.

The diagnostic value of the specificity, however, is plainly not affected by the phenomenon of group agglutination, since the action of minor agglutinins can be always easily eliminated by sufficient dilution. Thus if we possess a typhoid-immune serum which agglutinates the typhoid bacillus in dilutions of 1 to 10,000, the paratyphoid bacillus 1 to 1,000 and the colon bacillus 1 to 100 (as in the figure), we may still utilize this serum for the identification of suspected typhoid cultures, as, let us say, in the isolation of unknown bacteria from stools or urine, by using potent sera in dilutions as high or higher than 1 to 1,000, beyond which point the action of minor agglutinins is eliminated. The diagram illustrates our meaning in the hypothetical case of a typhoid-immune serum which agglutinates typhoid in dilutions of 1 to 10,000, paratyphoid bacilli 1 to 800, and colon bacilli 1 to 100. The relation of agglutination to biologic relationship is not a



DIAGRAMMATIC REPRESENTATION
OF GROUP AGGLUTINATION.

simple problem in that individual strains even of the same species may vary considerably in agglutination by the same serum. Smith and Reagh⁴¹ have studied particularly these conditions as they prevail in the colon, hog cholera and allied groups. They found that biologic relationship usually may be concluded from close agglutination affinities, and that minor biologic differences such as colony appearance, etc., do not exclude such affinities. On the other hand, closely related bacteria vegetating on mucous surfaces (different strains of diphtheria, dysentery, and colon bacilli) may vary considerably in their agglutinative characteristics, while invasive species show a greater homogeneity among their varieties or races. This brings in another important feature—that is, the modification in

⁴¹ Smith and Reagh. "Studies from the Rockefeller Institute," Vol. 1, 1904, p. 270.

agglutinative characteristics induced in bacteria when they become parasitic upon different hosts, and Smith and Reagh conclude that such changes of parasitic habitat may modify the agglutinative properties (probably by adaptation to the peculiar reactions of each host), some of them being weakened and others strengthened.

The animal species used for immunization indeed influences the quantity and nature of the produced agglutinin considerably. For instance, in Pfeiffer's ⁴² experiments, a dog, a chicken, and a rabbit were immunized with the same strain of cholera spirilla. The sera obtained from these animals agglutinated this and other strains of cholera spirilla in an entirely irregular manner—showing that the constitution of the agglutinins in each case was an absolutely different one in regard to the relative concentrations of "major" and "minor" constituents.

Castellani ⁴³ found that the immunization of an animal with two or more different species of bacteria results in the formation of agglutinins against all of these. Supposing, for instance, that species A and B are used for treatment, agglutinins against both A and B are formed in quantity, depending upon the intensity of the treatment in each case. Now, if to the serum so produced an emulsion of A is added, agglutinin A only will be removed, while agglutinin B will remain in the serum almost undiminished. An example of this is seen in the following protocol taken from Castellani's paper:

| Titre of the serum | Titre after absorption with <i>B. typhi</i> | Titre after absorption with <i>B. coli</i> "31" | Titre after absorption with <i>B. coli</i> and <i>B. typhi</i> |
|--|--|---|--|
| <i>B. typhi</i> 4,000 <i>B. coli</i> "31" 1,000 | <i>B. typhi</i> 0 <i>B. coli</i> "31" 1,000 > 300 | <i>B. typhi</i> 4,000 <i>B. coli</i> "31" 0 | <i>B. typhi</i> 0 <i>B. coli</i> "31" 0 |

In the preceding paragraphs, however, we have seen that immunization with a single organism, say *B. typhosus*, will induce the formation of agglutinins, not only for this species, but also of para or minor agglutinins for biologically similar strains as well. In such cases, as Castellani showed, absorption of the serum with the organism used for immunization takes out, not only the major agglutinins, but rather all of the agglutinins, major and minor. Conversely, however, absorption of such a serum with the species agglutinated by the minor agglutinins takes out these antibodies only, leaving the major substances intact. These relations are well illus-

⁴² Pfeiffer. Quoted from Paltauf in "Kolle u. Wassermann Handbuch," Vol. 4.

⁴³ Castellani. *Zeitschr. f. Hyg.*, Vol. 40, 1902.

trated by the two following protocols, also taken from Castellani's paper:

| Serum of rabbit No. 1 immunized with <i>B. typhi</i> only | | |
|---|---|--|
| Agglutination titre of serum | Titre after absorption with <i>B. typhi</i> | |
| <i>B. typhi</i> 5,000 <i>B. coli</i> 600 | <i>B. typhi</i> 0 <i>B. coli</i> 0 | |

| Serum of rabbit No. 7 immunized with <i>B. typhi</i> only | | |
|--|---------------------------------------|--|
| Agglutination titre of serum | After absorption with <i>B. typhi</i> | After absorption with <i>B. coli</i> |
| <i>B. typhi</i> 10,000 (heavy clumps) <i>B. coli</i> 800 | <i>B. typhi</i> 0 <i>B. coli</i> 0 | <i>B. typhi</i> 10,000 (small clumps) <i>B. coli</i> 0 |

Note: All of these protocols are taken from Castellani's communication, *loc. cit.*

These facts, variously confirmed, tend to corroborate the conception of the production of major and minor agglutinins outlined above.

It is of practical and theoretical importance to mention that complete absorption of specific agglutinin by a single exposure to homologous bacteria, however thickly emulsified, is not possible. It is always necessary to absorb repeatedly, and even then a minute trace of agglutinin may eventually remain. Eisenberg and Volk,^{44, 45} who have studied these conditions particularly, attribute this to the nature of the union of agglutino-gen with agglutinin, which they conceive as following the laws of mass action—this accounting for the persistence of a small "rest" of free agglutinin, even after repeated absorption by partial dissociation. The principle involved here is identical with that discussed in connection with antigen-antibody union in general.

It is not only in the sera of immunized animals, however, that agglutinins are found. Just as the other antibodies, antitoxins, and bactericidal sensitizers may be found in the blood of normal animals, so agglutinins for various bacteria may be normally present. These normal agglutinins do not in any respect, further than that of quantity, differ from the immune agglutinins and follow the same laws of specificity which have been described for the latter. It has been shown a number of times that such normal agglutinins are not pres-

⁴⁴ Eisenberg and Volk. *Zeitschr. f. Hyg.*, Vol. 40, 1902.

⁴⁵ Eisenberg. *Centralbl. f. Bakt.*, Vol. 34, 1903.

ent in the new-born animal, but are acquired later in life, possibly because of the absorption of bacterial products from the intestinal canal. It has been variously shown,⁴⁶ too, that living bacteria themselves may enter the lymphatics and the portal circulation from the intestine during apparently perfect health of the individual.

This subject is of interest, not only in connection with the agglutinins, but has bearing upon the existence of normal antibodies in general. Ruffer,⁴⁷ who has studied particularly the penetration of leukocytes and bacteria through the intestinal mucosa, demonstrated micro-organisms in the sub-mucous lymph nodes of normal rabbits, and Ribbert⁴⁸ and Bizzozero⁴⁹ have shown the presence of bacteria in apparently normal mesenteric lymph nodes. Adami and Nichols even claim that during health a certain number of living bacteria enter the portal circulation from the intestine, and from here may get into the systemic circulation, and are ordinarily destroyed by either leukocytes, liver lymphatic organs, or the kidneys.

It is thus not surprising that normal agglutinins should occur, and that they should be qualitatively identical with the so-called "immune" agglutinins, since they probably arise by a sort of spontaneous immunization through the intestinal canal. From the investigations of Ford especially we may conclude that the immune agglutinin may be regarded as merely a quantitative increase of the normal antibody, if this has been present before immunization. Ford⁵⁰ found that when an animal is treated with an agglutinating serum an anti-agglutinin may be obtained which neutralizes the action, not only of immune, but also of homologous, normal agglutinin.

An interpretation of the process of agglutination, according to the theory of Ehrlich, conceives it as a chemical union of agglutinin and bacteria (agglutino-gen). The agglutinin is regarded as consisting of two atom complexes, one the "haptophore," having affinity for the bacterial protein, and concerned with the union, the other the "ergophore" or "zymophore," by means of which the actual agglutination is brought about after the union has taken place. Unlike the antibody concerned in the processes of hemolysis or bacteriolysis, the agglutinins are not dependent in their action upon the coöperation of alexin, and the agglutination power of a serum is therefore not destroyed by inactivation or heating to 56° C., as is the case with the former. Although the accurate point of thermal destruction varies with different agglutinins (the agglutinins for the *Bacillus pestis* and a few other bacilli are said to be destroyed at

⁴⁶ Adami. *Jour. Am. Med. Assoc.*, Dec., 1899.

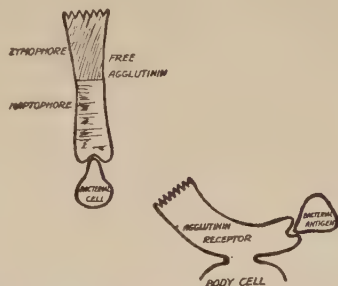
⁴⁷ Ruffer. *Brit. Med. Journal*, 2, 1890.

⁴⁸ Ribbert. *Deutsche med. Woch.*, 1885.

⁴⁹ Bizzozero. *Centralbl. f. d. Med. Wiss.*, Vol. 23, 1885, p. 49. Quoted from Adami.

⁵⁰ Ford. *Zeitschr. f. Hyg.*, Vol. 40, 1902.

56° C.), as a rule agglutinins will not disappear from serum until the temperature is raised to between 70° and 80° C. Once destroyed, however, no reactivation takes place upon the addition of fresh normal serum. Ehrlich, for this reason, has conceived the structure of agglutinins as "*Haptines* of the Second Order," in which he supposes that the zymophore and the haptophore groups are inseparably connected, and in which we could assume an alteration of the less stable zymophore group without interference with the functional integrity of the haptophore group. Such an altered agglutinin could be spoken of as "agglutinoid," and this could become united with a bacterial cell without inducing agglutination, but, by its union, prevent subsequent combination of the cell with unaltered agglutinin. This process of "agglutinin Verstopfung" has been held responsible for the failure of agglutination when bacteria that have been in contact with heated serum are subsequently exposed to the action of actively agglutinating serum. It is assumed that the agglutinoids which were present in the heated serum have occupied the bacterial receptors and have thereby prevented the union of these with the agglutinins later added.



DIAGRAMMATIC REPRESENTATION OF
EHRlich's CONCEPTION OF THE
STRUCTURE OF AGGLUTININ.

The work of Eisenberg and Volk⁵¹ has gone very thoroughly into these conditions and forms the strongest bulwark of this point of view. These workers showed that bacteria thus exposed have not only become less sensitive to agglutinins, but have, at the same time, lost much of their power to absorb agglutinins when compared with normal bacteria. The same loss of agglutinating power which is observable in heated agglutinating serum is evident to a lesser extent in serum which has been preserved at room temperature. Eisenberg and Volk have shown that such serum, in addition to a loss of its total agglutinin content, loses the power to agglutinate in high concentrations. Thus a serum which has been preserved in this way will no longer agglutinate bacteria in concentrations of 1 to 20, 1 to 40, or even 1 to 100, but will agglutinate as before in higher dilutions. This is taken to mean that the agglutinoids formed during the period of standing possess a higher affinity for the bacterial antigen than do the true unaltered agglutinins. Since these so-called "proagglutinoids" are relatively small in amount, their action is masked when considerable dilution has sufficiently diminished their quantity, in proportion to the more plentiful un-

⁵¹ Eisenberg and Volk. *Zeitschr. f. Hyg.*, Vol. 40, 1902.

altered agglutinins. In support of this assumption it has been further shown that sera which have been rendered inhibitory by either of the methods named can be deprived of their inhibiting characteristics by absorption with homologous bacteria. Together these observations constitute a strong argument in favor of the agglutinoid theory.

In practical experience the existence of such an inhibition zone is of great importance, since freshly taken sera will occasionally show this failure of agglutination in concentration, while strong agglutination follows when the dilution is increased. In clinical tests, as in the Widal reaction for the diagnosis of typhoid fever, we not infrequently encounter sera which will give no agglutination in dilutions of 1 to 20 and even 1 to 40, and the reaction would therefore be regarded as negative unless the possibility of the proagglutinoid zone were recognized and further dilutions carried out.

While there is no question of the accuracy of the experimental data cited in the preceding paragraphs, the interpretation of the phenomena on the basis of Ehrlich's haptine conception has not been universally accepted.

The fact that the agglutinins lose their agglutinating power after heating, while retaining their power to prevent the subsequent agglutination of the bacteria, may be more simply explained in analogy with the observation of Porges on the influence of heated serum upon the agglutination of mastic suspensions. He found that unheated serum will flocculate such suspensions, while heated serum of the same concentration will prevent the flocculation, acting probably as a protective colloid (see chapter on Colloids). In the same way the heated agglutinating serum may prevent subsequent flocculation by a similar protective action. We suggest this as a possible explanation of the proagglutinoid phenomenon, although of course it is a mere conjecture as opposed to the painstaking experiments of Eisenberg and Volk. It has the advantages of simplicity, but does not, it is true, explain the apparent specific absorption of the agglutinin-inhibiting properties out of heated sera with homologous bacteria, as claimed by these authors as well as Kraus and Joachim. The similarity of the proagglutinoid phenomenon to the inhibition zones occurring in the flocculation of colloids will be referred to in a subsequent paragraph.

In describing the investigations which led to the discovery of the mechanism of the lytic phenomenon, in the chapter on Cytolysis, we mentioned that Bordet and others had noticed the frequent agglutination of red blood cells in the sera of animals treated with such cells after the hemolytic property had been destroyed by heating to 56° C. Such hemagglutination is a phenomenon entirely analogous to the agglutination of bacteria by serum, and hemagglutinins regularly result when an animal is systematically treated

with the red blood cells of another species. Like the bacterial agglutinins, the hemagglutinins are relatively thermostable and are best observed, therefore, after the sera are inactivated. Otherwise rapid hemolysis will often obscure the agglutination. Like other agglutinins the hemagglutinins thus produced are specific, acting only upon that variety of cells which are used in their production. Moreover, certain sera may normally contain hemagglutinins for the blood cells of animals of another species. An illustration of this is the hemolytic and hemagglutinating property of normal goat serum for rabbit cells—but there are many other examples which might be cited. Such normal hemolytic and hemagglutinating properties for the cells of other animals usually render the sera toxic for these animals, and some observers have attributed the toxicity to this agglutinating action, believing that the clumped erythrocytes form emboli around which clotting is initiated. The writer's own investigations, however, seem to show that this is not the case, since the toxicity of such sera is completely removed after they have been heated, in spite of the fact that the hemagglutinative property remains unchanged.

In discussing hemolysins, also, we called attention to the observation that the sera of animals may develop the property of hemolyzing blood cells of other individuals of the same species when immunized with such cells, and that on occasion such isolysins may be normally present.

Analogous to iso-lysins, iso-agglutinins also have been observed. They were described first in human blood in 1900, independently, by Landsteiner,⁵² and by Shattock. As the result of the work of a number of men, in particular that of Landsteiner, of Ascoli,⁵³ and of Descatello and Sturlii,⁵⁴ it became evident that all human blood fell into four sharply separated and, for the individual, permanent groups, according to the way in which they interagglutinate. The members of the first group have blood cells which are not agglutinated by the serum of any human blood. The serum of the members of this group agglutinates the blood cells of persons belonging to any of the other three groups. This group constitutes between 40 to 50 per cent. of all human beings. Members of the second group have blood serum which agglutinates the cells of persons belonging to the third and fourth groups; while the cells of the second group are agglutinable by the serum of the first and third groups. The third is the reciprocal of the second, and the serum of the third group agglutinates the cells of the second and fourth groups; while the cells of the third group are agglutinated only by the serum of the first and second groups. The fourth group (which is very rare) is the recipro-

⁵² Landsteiner and Richter. *Zeitschr. f. Med.*, 3, 1902.

⁵³ Ascoli. *Münch. med. Woch.*, 1901.

⁵⁴ Descatello and Sturlii. *Münch. med. Woch.*, 1902, 49, p. 1090.

TABLE FOR ISO-AGGLUTININS *

Sera

| | | I | | | | II | | | III | | IV |
|-----|----|---|---|---|---|----|---|---|-----|---|----|
| | | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 |
| I | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| | 2 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| | 3 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| | 4 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| II | 5 | + | + | + | + | 0 | 0 | 0 | + | + | 0 |
| | 6 | + | + | + | + | 0 | 0 | 0 | + | + | 0 |
| | 7 | + | + | + | + | 0 | 0 | 0 | + | + | 0 |
| III | 8 | + | + | + | + | + | + | + | 0 | 0 | 0 |
| | 9 | + | + | + | + | + | + | + | 0 | 0 | 0 |
| IV | 10 | + | + | + | + | + | + | + | + | + | 0 |

* For this table as well as for much direct information concerning the iso-agglutinins and isolyisin we are indebted to Dr. Ottenberg of this laboratory.

cal of the first, the serum having no agglutinin, the cells being agglutinated by the serum of any other group. (See table.) It is evident from examining this grouping that the phenomena can be explained (as Landsteiner has suggested) if it is assumed that there are two agglutinins (α and β) and two corresponding agglutinogens present in the red cells (A and B). The blood of the first group possesses both agglutinins, but no agglutinogens, the blood of the second group possesses agglutinin α , agglutininogen B, the blood of the third group possesses agglutinin β , agglutininogen A, the blood of the fourth group possesses no agglutinin but both agglutinogens.

These agglutinins are present in weak dilution only, being generally active in dilutions only of 1-15 to 1-30. They are separately absorbed from the serum by the suitable red cells (Hektoen).⁵⁵ Ot-

⁵⁵ Hektoen. *Jour. of Inf. Dis.*, 1907, p. 297.

tenberg noticed that they were inherited, and this was also shown in 1908 and in 1910 by von Dungern and Hirschfeld,⁵⁶ who further found that this inheritance followed the Mendelian law strictly. The agglutinogens are the unit characters. The agglutinogens apparently are present at an earlier embryonic stage than the agglutinins. On account of their hereditary nature and permanence for the individual the iso-agglutinins may possibly be of medicolegal value. They may also be of some practical importance in selecting donors for blood transfusion, as phagocytosis of red blood cells in the circulation after transfusion, first described by Hopkins, was proved by Ottenberg to occur only when the patient's serum was agglutinative toward the donor's red cells, and several such transfusions have had fatal results. Similar iso-agglutinins have been observed in the blood of lower animals, in horses (Klein,⁵⁷ 1902); rabbits (Boycott and Douglas,⁵⁸ 1910); cats (Ingebrigtsen); dogs, rats, and steers (Ottenberg).⁵⁹ The iso-agglutinins have been developed in dogs (von Dungern and Hirschfeld).⁶⁰ In most of the lower animals they have occurred with peculiar irregularity, indicating probably the presence of, not two, but of a larger number of agglutinins and agglutinogens. In steers, however, they fall into simple groups, indicating the presence of only one agglutinin and agglutinogen. In many animals the agglutinins are entirely latent, and the biochemical differences represented by the agglutinogens are present in the red cells, and the correct agglutinin is developed by the animal only when it is immunized with blood whose cells contain agglutinin not present in the animal's own blood cells.

The fundamental principle underlying all the Ehrlich hypotheses is the conception that these reactions take place as do purely chemical reactions, following the law of multiple proportions. Such reasoning has often necessitated the assumption of differences of affinity which, critically examined, are really *ex post facto* explanations, forcing the phenomena to conform with the theory. As a matter of fact, the bodies which participate in the antibody-antigen reactions are probably of the nature of the substances which are spoken of as colloids, and it is therefore more than likely that the quantitative and other relations governing the union of these reagents should be analogous to those governing colloidal reactions in general. The reaction of agglutination, like that of precipitation, has lent itself particularly to the study of the principles of the union from this point of view, and the first and fundamental progress made in this direction is found in the work of Bordet.

⁵⁶ Von Dungern and Hirschfeld. *Zeitschr. f. Imm.*, 4, 1909-1910, p. 53; also *ibid.*, 1910, p. 284.

⁵⁷ Klein. *Wien. kl. Woch.*, 1902, p. 413.

⁵⁸ Boycott and Douglas. *Jour. of Path. and Bact.*, Jan., 1910.

⁵⁹ Epstein and Ottenberg. *Tr. N. Y. Path. Soc.*, 1908.

⁶⁰ Von Dungern and Hirschfeld. *Zeitschr. f. Imm.*, 1909, 1910, p. 531.

Bordet⁶¹ compared the formation of precipitates in bacterial emulsions to the precipitation of such inorganic emulsions as clay in distilled water, and noted that the precipitation of homogeneous emulsions of such substances is "often controlled by such insignificant causes as the presence of salts." Applying this analogy to the agglutination of bacteria, he performed the following experiment: Cholera spirilla, emulsified in salt solution, were treated with homologous immune serum and, after agglutination had taken place, the bacteria were thrown to the bottom by centrifugation and divided into two parts. One part was again suspended in salt solution, and the other was washed, and then suspended in distilled water. The bacteria in the tube of salt solution rapidly agglutinated, while those in the distilled water, after thorough shaking, remained indefinitely suspended in an even emulsion. If, however, to these unagglutinated bacteria a small amount of pure sodium chlorid was added agglutination occurred.

The conclusions that can justly be drawn from this experiment are, first, that the bacteria could not agglutinate, even though they had been bound to agglutinin, when salt was removed from the environment, and, second, that the addition of salt to such emulsions brought about immediate agglutination. The same principle can be demonstrated in other ways. If, for instance, a bacterial emulsion is rendered free of salt by dialysis, and this is added to an agglutinating serum similarly dialyzed, no agglutination occurs. The suspension may remain evenly clouded indefinitely unless salt is added. As soon as a little salt is added, however, perfect agglutination occurs. To this technique the very obvious criticism may be applied that perhaps the absence of salt has precipitated the agglutinins, which, as we know, are precipitated with globulin, which is insoluble in the absence of salt. However, this source of error is excluded by the first experiment cited, and, moreover, it can be shown by the last experiment that, even though the bacteria are not agglutinated in the salt-free serum, they have nevertheless absorbed agglutinin. For, if such a salt-free mixture is centrifugalized, the bacteria washed and suspended in distilled water, and salt is then added, agglutination occurs. The supernatant fluid of the original suspension, furthermore, can be shown to have been deprived of agglutinins by suitable experiment.

These facts, first observed by Bordet, and further elaborated by the studies of Joos,⁶² Friedberger,⁶³ and others, have been interpreted in different ways. Joos claims that there is a chemical union between the bacteria and the salt, and bases this upon the observation that the salt added to a salt-free mixture cannot be demonstrated in

⁶¹ Bordet. *Ann. de l'Inst. Pasteur*, 1896, 1899.

⁶² Joos. *Centralbl. f. Bakt.*, 1, Vol. 33, 1903.

⁶³ Friedberger. *Berl. kl. Woch.*, 1902; *Centralbl. f. Bakt.*, 1, Vol. 30.

the supernatant fluid after agglutination has taken place. His observations in this respect have not found confirmation at the hands of Friedberger and other workers, and it is generally agreed to-day that the rôle of the salts is, as Bordet first assumed it to be, a purely physical one. Bordet's opinion is often spoken of as the "two phase" theory, in that he conceives the process of agglutination to consist of two distinct occurrences, first, an absorption of the agglutinin by the bacteria, and, second, an agglutination of the new complex by the salt. It is not the agglutinin which causes agglutination, but by union with the agglutinin forms a complex which is altered in "colloidal stability," and therefore is flocculable by the electrolyte.

The opinion of Bordet becomes clearer as we consider the conditions governing the flocculation of colloids in general. Without wishing to enter in this place into detail regarding the nature of colloidal suspensions, it nevertheless seems necessary in order to do justice to this phase of the question to recall briefly the conditions governing such flocculation. The so-called colloidal solutions are not true solutions as the term is applied to dissociable substances, but are looked upon as consisting of small particles in suspension. The particles are similarly charged, as can be demonstrated by their wandering when subjected to an electric current, and it is supposed that it is this fact of similarity of charge which, in the "sol" state, permits them to remain in suspension. For the similarity of the charges of the individual particles prevents their mutual approximation.

The state of suspension of these substances, then, represents a delicately balanced equilibrium between the two forces of electrical repulsion and of surface tension, an equilibrium which may be disturbed by the action of a number of factors. Thus, studies on inorganic colloids have shown, long before these considerations were applied to the explanation of serum reactions, that the stability of these suspensions could be disturbed both by electrolytes and by the addition of other colloids. Thus they may be precipitated by various salts, acids, and bases and, as Schultze⁶⁴ has shown, they react with that ion of the electrolyte which carries an opposite charge to that of the colloidal particles. For, although the colloidal units are similarly charged, this may be either negative or positive, according to the nature of the particular substance. In the case of the so-called amphoteric colloids reaction may take place, according to Pauli,⁶⁵ with both ions of the electrolyte.

The probable mechanism of the process is postulated by Pauli in describing the precipitation of a colloidal metal by salts, acids, or bases in the following way:

⁶⁴ Schultze. *Journ. f. prakt. Chem.*, 25, 1882, and 27, 1883.

⁶⁵ Pauli. "Hofmeister's Beiträge," 1905, and "Physical Chemistry in Medicine," Wiley & Son, N. Y., 1907.

"The introduction of such electrolytes into a colloidal suspension is of course accompanied by a certain amount of dissociation. In consequence the weakly charged particles of the colloid collect about the ions of opposite charge until a sufficient accumulation of the particles leads to an electrical neutralization of the ion, and the aggregation, if of sufficient size, will sink to the bottom, forming a precipitate."

In regard to the mutual influences exerted upon each other when two colloids are mixed, it has been shown by Biltz, Hardy, and many other observers that oppositely charged colloids precipitate each other, though this is not an absolute rule, as experiments by Professor Stewart Young, of Stanford, have shown. Thus colloidal gold and platinum will be precipitated by such colloids as ferric oxid or aluminium oxid. When such a precipitating colloid is added to another oppositely charged suspension in quantities too small to bring about flocculation, moreover, the addition of a quantity of salt, likewise too small to precipitate alone, will in many cases bring about the flocculation.

These and other phenomena of colloidal reaction have found close analogy in antibody-antigen studies, and have given support to the interpretation of the latter in the sense of Bordet.

To return to the consideration of bacterial agglutination, we have spoken of the dependence of the reaction upon the presence of salts, and have seen that the researches of Friedberger and others have refuted the assumption that the action of the salt in bringing about agglutination depends upon chemical union of the salt with the bacteria. It is probable, therefore, that here, as in other colloidal precipitations, the function of the salt is to be regarded purely as an electrophysical phenomenon.

The analogy becomes still closer when we consider the researches of Bechold,⁶⁶ Neisser and Friedemann,⁶⁷ Sears and Jameson,⁶⁸ and others, which have shown that bacteria in suspension are to be compared very closely with true colloidal suspensions in that the bacterial cells carry a definite and uniform electrical charge.

Bacteria in salt solution emulsion, for instance, wander to the anode, thus giving evidence of their carrying a negative charge. This charge may be altered by adding to the emulsions definite concentrations of acids or bases, a reversal of the charge taking place under the influence of NaOH or other hydroxids. Just how this is brought about is by no means clear, but it is not impossible that there is a selective absorption of OH ions by the bacteria, which therefore take on the charge of the ion.

⁶⁶ Bechold. *Zeitschr. f. physik. Chem.*, 48, 1904.

⁶⁷ Neisser and Friedemann. *Münch. med. Woch.*, Vol. 51, pp. 465, 827, 1904.

⁶⁸ Sears and Jameson. "Thesis for M. A. Stanford University," 1912.

However this may be, and we must admit that explanations of these phenomena are as yet largely speculative, a fact which interests us particularly in connection with the phenomena under discussion at present is the influence exerted upon the charge of bacteria by exposure to the influence of serum. Bechold,⁶⁹ as well as Neisser and Friedemann,⁷⁰ assert that bacteria which have absorbed agglutinin no longer wander to the anode, but act as though they had been deprived of electrical charge, and precipitate between the electrodes.

Bechold has suggested, for this reason, that it may be possible that bacteria in the normal condition are protected from the action of the electrolyte by a membrane or coating of protoplasm which acts as a protective colloid. The absorption of agglutinin may alter this in such a way that they become amenable to the flocculating effects of the salt ions. In keeping with such an opinion is the well-known observation of the inagglutinability of capsulated organisms, which, as Porges⁷¹ has shown, become agglutinable as soon as the capsules have been destroyed by heating in a weak acid.

That the absorption of agglutinin indeed alters the electric stability of the emulsified bacteria further appears from the fact that "agglutinin" bacteria⁷² are agglutinated by concentrations of salts which are too slight to affect the normal micro-organisms. In this respect there is close similarity between the flocculation of agglutinin-bacteria and such emulsions as kaolin and mastic, whereas bacteria without agglutinin require much higher concentrations of the salts to produce like effects. The absorption of agglutinin may have removed a factor which protected the bacteria against the influence of the salt. On the other hand, it is equally just to assume—and this is more likely and corresponds with Bordet's views—that the absorption of agglutinin has "sensitized" the bacteria to the action of the electrolyte. The experimental facts upon which the above statements are formulated are largely found in the important papers of Neisser and Friedemann—whose work brought out, likewise, interesting analogies of the colloidal precipitations with the phenomenon which we have described above as the proagglutinoid zone.

In regard to the greater amenability of agglutinin bacteria to flocculation by electrolytes, the following protocol, adapted from the work of these authors, will explain itself. They were tabulated from experiments in which different quantities of normal $\frac{1}{1}$ solution of various salts were added, on the one hand to emulsions of unal-

⁶⁹ Bechold. "Die Kolloide in der Biologie u. Medizin," Steinkopf, Dresden, 1912.

⁷⁰ Neisser and Friedemann. *Münch. med. Woch.*, Vol. 51, 1904, pp. 465 and 827.

⁷¹ Porges. *Ztschr. f. exp. Path. u. Therapie*, 1905.

⁷² "Agglutinin" bacteria—bacteria which have absorbed specific agglutinin.

tered bacteria, and, on the other, to bacteria which had absorbed agglutinin. It is seen that, with some salts, agglutination of the unaltered bacteria did not occur at all, whereas agglutination was brought about in the treated bacteria with comparatively small amounts; in other cases the difference is a quantitative one only:

Protocol constructed from the tables of Neisser and Friedemann, loc. cit.

| $\frac{n}{1}$ solution of salt | Quantity of salt sol. which brought about agglutination of 1 c. c. of normal bacteria in emulsion. 0 = no agglutination by the salt solution | Quantity of salt sol. which agglutinated 1 c. c. of agglutinin bacteria in emulsion |
|---|---|---|
| NaCl..... | 0 | .025 |
| NaNO ₃ | 0 | .025 |
| Na ₂ SO ₄ | 0 | .025 |
| RbI..... | 0 | .025 |
| MgSO ₄ | 0 | .0025 |
| ZnSO ₄ | .01 | .001 |
| CaCl ₂ | 0 | .005 |
| BaCl ₂ | 0 | .005 |
| Cd(NO ₃) ₂ | .01 | .001 |
| CuSO ₄ | .0025 | .0001 |
| CuCl ₂ | .0025 | .0005 |
| Pb(NO ₃) ₂ | .0025 | .0001 |
| HgCl ₂ | .0025 | .0005 |

The analogy between the experiment tabulated in the preceding protocol and the following from the work of the same writers is self-evident. Just as the absorption of agglutinin by bacteria rendered these more amenable to precipitation by salts, so the addition of minute quantities of gelatin to mastic emulsions had a similar sensitizing effect upon these.

| NaCl 10% solution | 1 c. c. mastic (1-10 original emulsion) diluted to 3 c. c. | 1 c. c. mastic + .0001 c. c. of a 2% gel. solution, the whole diluted to 3 c. c. |
|----------------------|--|--|
| 1.0 | +++ | +++ |
| 0.5 | 0 | +++ |
| 0.25 | 0 | +++ |
| 0.125 | 0 | +++ |
| 0.05 | 0 | 0 |
| 0.025 | 0 | 0 |

Finally, one of the most important analogies yielded by the work of the above investigators is illustrated in the following protocol:

| Colloidal iron hydroxid | Precipitation of mastic emulsion 1 c. c. |
|----------------------------|---|
| 1. | 0 |
| 0.5 | 0 |
| 0.25 | 0 |
| 0.1 | ++ |
| 0.05 | ++++ |
| 0.025 | ++++ |
| 0.01 | ++++ |
| 0.005 | ++++ |
| 0.0025 | ++ |
| 0.001 | 0 |

Here we have an inhibition zone in the tubes containing the highest concentrations, accurately analogous to the previously discussed *proagglutinoid* zone. It is a phenomenon similar also to the inhibition zones noticed in precipitin reactions and observed, though by a different technique, in bacteriolytic phenomena discussed in another place in connection with the Neisser-Wechsberg notion of complement-deviation or "Komplement Ablenkung." It seems to be a universal fact governing the union of colloidal substances, that definite quantitative proportions must be maintained in order to lead to reaction, this being, possibly, explicable on the basis that actual union can take place only after disturbance of the electrical balance which keeps the particles apart. These reactions will be found more accurately discussed in another place. Whatever the mechanisms may be, however, these and similar experiments have seemed to render unnecessary and unlikely the assumption of proagglutinoids, proprecipitoids, etc., to explain the inhibition zones so frequently observed in all reactions of this kind.

A peculiar observation, which does not coincide with the above interpretation of these phenomena, and the significance of which is indeed doubtful, is one which Friedberger⁷³ made in researches in which he confirmed the work of Bordet on the absence of agglutination in a salt-free environment. He found that not only the addition of various salts would bring about agglutination under such conditions, but that organic substances such as dextrose and asparagin could be substituted for salts and had similar agglutinating effect—although higher concentrations of these than of the salts were required. Were these substances at all dissociable it might be possible that they acted by a mechanism identical with that of the salts—but since such substances as dextrose either do not dissociate at all or do so to an infinitesimal degree only there does not seem any possibility of reconciling these results with Bordet's theory.

⁷³ Friedberger. *Centralbl. f. Bakt.*, 30, 1901.

It is difficult to explain Friedberger's results. Possible impurity of his preparations and the presence of traces of electrolyte seem to be excluded by the fact that he was quite conscious of this possibility of error and used only substances which yielded no ash on combustion.

It may be that the results of Friedberger in which glucose and asparagin were used may have brought about agglutination by an entirely different mechanism from that which we are discussing and form no analogy to this.

In one of the preceding paragraphs we have mentioned the phenomenon spoken of as "acid agglutination." By this is meant the spontaneous clumping, not only of bacteria, but of small particles of any kind, in suspension, in the presence of certain concentrations of acid. Michaelis,⁷⁴ Beniasch,⁷⁵ and others who have studied this phenomenon in detail have come to the conclusion that it is the concentration of the hydrogen ions which is responsible for the agglutination. This explanation is also applicable to the agglutination often observed about the anode when bacteria are subjected in suspension to the action of a direct current. In such experiments the organisms after concentrating at this electrode often flocculate, and it is here, of course, that hydrogen ions are present in the greatest concentration. How this takes place is problematical, but the reasoning of Pauli, if applied to this, would favor the assumption that the weakly charged bacteria group themselves about the ions and, when a sufficiently large aggregation has formed, fall to the bottom as precipitate. This phenomenon of acid agglutination is of course entirely different in nature from the specific serum agglutination which we are discussing. Nevertheless, Schidorsky and Reim,⁷⁶ Jaffe,⁷⁷ and others have attempted to apply acid agglutination to the isolation and differentiation of bacteria, on the conception that different species are agglutinated by varying concentrations of hydrogen ions. The former investigators, even, claim to have been successful in isolating typhoid bacilli from the stools by this method in that the typhoid bacillus was agglutinated by concentrations of acid which had no effect upon the *Bacillus coli*. Sears⁷⁸ has gone over this work carefully, and, while he has obtained results which bear out the contention that the agglutination is probably due to the concentration of the H ions, his experiments have revealed an irregularity in the behavior of bacteria of the same species in acid solutions and an overlapping of those of one species with those of another. Therefore the use of acid agglutination for differential purposes

⁷⁴ Michaelis. *Folia Serol.*, 7, p. 1010, and *Deutsche med. Woch.*, 37, 969.

⁷⁵ Beniasch. *Zeitschr. f. Imm.*, Vol. 12, 1912.

⁷⁶ Schidorsky and Reim. *Deutsche med. Woch.*, Vol. 38, p. 1125.

⁷⁷ Jaffe. *Arch. f. Hyg.*, Vol. 76.

⁷⁸ Sears. *Proc. Soc. of Exp. Biol. and Med.*, 1913.

seems to us entirely hopeless. And indeed it would be surprising if any such distinctive and regular reaction differences between simple bacterial cells, after all chemically and physically so essentially alike, could be found.

CHAPTER X

THE PHENOMENON OF PRECIPITATION (Precipitins)

THE establishment of the agglutinin reaction as a constant and specific serum-phenomenon by the work of Gruber and Durham led immediately to assiduous investigation of the many problems suggested by it, and among them, as we have seen, the question of the nature of the agglutinin. It was found that agglutinins could be produced, not only by the injection of whole bacteria, but equally as well by treatment with dissolved bacterial extracts or with filtrates from old broth cultures. This naturally led to the thought that there might be a definite reaction if such extracts (instead of the bacteria themselves) were added to agglutinating sera *in vitro*. Rudolf Kraus¹ was the first to perform this very logical experiment. He was working with broth filtrates of *Bacillus pestis* and of the cholera spirillum, and found that when he mixed the perfectly clear filtrates of such cultures with their respective antisera the mixtures would at first become turbid and finally show a light flocculent precipitate. He named the reaction the "precipitin reaction" and, in analogy to agglutinins, spoke of the bodies in the serum which caused the precipitation as "precipitins." The reaction was found, like that of agglutination, to be specific; the cholera serum gave no precipitate with the plague extract and *vice versa*, and Kraus, after extending his observations to other bacteria, pointed out the practical diagnostic possibilities of his discovery.

Though Kraus' first observations were made entirely with bacterial culture filtrates and antibacterial sera, it was soon discovered that his results were merely isolated instances of a broad biological law, and that specific precipitins were produced whenever animals were treated with injections of any kind of foreign protein. Thus Tschistovitch,² in 1899, found that the blood serum of rabbits immunized with eel-serum gave specific precipitates when mixed with eel-serum, and Bordet³ obtained analogous results by treating rabbits with defibrinated chicken blood and with milk. Thus rapidly

¹ R. Kraus. *Wien. klin. Woch.*, No. 32, 1897.

² Tschistovitch. *Ann. de l'Inst. Past.*, 13, 1899.

³ Bordet. *Ann. de l'Inst. Past.*, Vol. 13, 1899, pp. 225-273.

the discovery of Kraus was developed into the generalization that the sera of animals that have been treated with foreign proteins of any kind—bacterial, animal, or vegetable—will develop the property of causing precipitates when mixed with clear solutions of the respective antigens.

The substances which, after injection into the animal body, lead to the formation of precipitating antibodies are spoken of in the language of immunology as "precipitinogen." In the case of bacteria it has been shown that, while the injection of the whole bacterial cell—dead or alive—will lead to precipitin formation, bacterial extracts produced in a variety of ways will lead to the same result. Such precipitinogen extracts can be obtained by allowing the bacteria to grow in flasks of slightly alkaline bouillon, keeping them in the incubator for from three weeks to three months, and then filtering them through Berkefeldt candles. Again, useful extracts can be more rapidly produced by growing large quantities of bacilli on agar, emulsifying in salt solution, and shaking in any one of the ordinary types of shaking machine for 48 hours or longer. On filtering an extract is obtained which will form precipitates with homologous immune serum, or will incite precipitins when injected into animals. In fact, any one of the customary vigorous methods of extracting bacterial or other cells will yield precipitinogen. A relatively purified precipitinogen in the form of a dry, water-soluble powder has been obtained by Pick by the precipitation of culture filtrates with alcohol.

Regarding the chemical nature of the precipitin-inducing substances, or precipitinogens, the same problems have arisen which have been discussed in connection with antigens in general. We may say that all soluble native proteins possess precipitin-inducing properties. Yet this does not sufficiently define the term, since many observations have been published which show that physically and chemically altered proteins may still induce specific precipitins; a few investigators, furthermore, have claimed that they have produced non-protein precipitinogen by various methods of breaking up the molecule of the original antigen. In the section on agglutination we have seen that moderate heating (56-65° C.) rather increases than decreases the agglutinin characteristics of bacteria, and it is equally true that such heated bacteria or bacterial extracts may induce precipitins. However, regarding the action of higher degrees of heat (boiling) upon precipitinogens in general we will have more to say in another place.

Of more immediate, indeed of fundamental, importance is the problem of a non-protein antigen. The most important claims in this regard have been made by Pick,⁴ Obermeyer and Pick,⁵ and by

⁴ Pick. "Hofmeister's Beiträge," Vol. 1, 1901.

⁵ Obermeyer and Pick. *Wien. klin. Woch.*, 1904, p. 265.

Jacoby.^{6 7} Jacoby, working with a vegetable antigen, ricin, found that by trypsin digestion he could obtain a substance which still retained antigenic properties, but no longer gave any of the protein reactions. Obermeyer and Pick, by the same method, claim that they have produced a non-protein precipitinogen from egg albumen. On the other hand, others have had negative results, and Kraus⁸ himself, after reviewing the evidence on both sides, comes to the conclusion that available data do not justify us in separating the antigenic properties from the protein molecule. In unpublished experiments which the writer carried on in the laboratory of Professor Friedemann in Berlin also attempts to produce a non-protein precipitinogen from horse serum by bacterial putrefaction were entirely negative. The putrefaction of the serum, though carried out in dialyzing bags for the removal of diffusible products, was extremely slow, and when finally the Biuret reaction disappeared the serum was no longer precipitable by potent antisera. However, the flaw in these experiments is that the true test of the presence of precipitinogen is not the precipitable character of the solution in question, since actual precipitation is dependent, as we shall see, upon many modifying secondary factors, but rather the ability of the substance to induce precipitins in treated animals.

The fact that Nicolle,⁹ and later Pick,¹⁰ were unable to obtain alcohol-soluble substances from bacteria and bacterial extracts which were still precipitable might also be taken to point toward the non-protein character of the precipitinogens, suggesting that these substances may be of a lipoidal nature. However, as Landsteiner¹¹ points out, mere solubility in organic solvents can no longer be taken as a proof of lipoidal character, since it is more than probable that non-lipoidal substances may go into alcoholic and other organic solution when lipoids, such as lecithin, are present. Thus Müller¹² found that the antigen of typhoid bacilli was soluble in chloroform in the presence of old preparations of lecithin. Pick and Schwartz,¹³ who had previously studied similar antigen solubilities in the presence both of lecithin and of other organ lipoids, suggest that possibly such solutions represent lipid-protein combinations—colloidal “solutions”—which permit the presence of protein mechanically or chemically united to the lipid in the organic solvents—alcohol, chloroform, etc. Here, too, then there is no evidence for the existence of non-protein precipitinogen.

⁶ Jacoby. “Hofmeister’s Beiträge,” Vol. 1, 1901.

⁷ Oppenheimer. “Hofmeister’s Beiträge,” Vol. 4, 1904, p. 259.

⁸ Kraus in “Kolle u. Wassermann Handbuch,” Vol. 4, p. 605.

⁹ Nicolle. *Ann. de l’Inst. Past.*, 12, 1898.

¹⁰ Pick. “Hofmeister’s Beiträge,” Vol. 1, 1901.

¹¹ Landsteiner. “Weichhardt’s Jahresbericht,” Vol. 6, 1910, p. 214.

¹² Müller. *Zeitschr. f. Imm.*, Vol. 5, 1910.

¹³ Pick and Schwartz. *Biochem. Zeitschr.*, Vol. 15, 1909.

Of importance in connection with the problem of the nature of precipitinogen, also, is the claim of Myers,¹⁴ that specific precipitins may be produced in rabbits by treatment with Witte peptone, a substance complex in constitution, but consisting largely of albumoses. This observation has failed of confirmation in the hands of Obermeyer and Pick, Michaelis,¹⁵ Norris,¹⁶ and others, and cannot, therefore, be accepted as an established fact.

Whichever method of precipitinogen production is used bacterial precipitins appear in the serum of the immunized animal only after careful and continued immunization, usually later than the demonstrable appearance of the bactericidal or agglutinating properties of the serum. The most convenient material for such immunization consists of salt solution emulsions of agar cultures, killed at 60° to 70° C. These may be injected subcutaneously, intraperitoneally, or intravenously, the last method leading to the most satisfactory and rapid results and, therefore, best employed unless great inherent toxicity of the particular bacteria contraindicates. When rabbits are used it is generally necessary to inject 3, 4, or 5 times at 5 or 6-day intervals, and to bleed the animals on the 8th or 9th day after the last injection.

The bacterial precipitins so produced are, as we have said above, specific—but, again, specificity, as in the case of agglutinins, is limited by the so-called “group reactions.” In the chapter dealing with agglutination we have seen that the serum of a typhoid-immune animal which agglutinates typhoid bacilli strongly will also agglutinate, though far less powerfully, paratyphoid bacilli and, in some cases, even colon bacilli, this appearance of “minor” agglutinins being probably due to a close group relationship of these bacteria to the typhoid bacillus. In the case of bacterial precipitins the same thing is true, and has been made the subject of special studies by Zupnik,¹⁷ Kraus,¹⁸ Norris,¹⁹ and others. As in the case of agglutination, however, this fact does not in any way interfere with the practical value of the specificity of the reaction because elimination of the secondary group reactions, which in agglutination is obtained by dilution of the antiserum, can here be obtained, as Kraus points out, by diminishing the quantity of the undiluted precipitating serum added to the bacterial filtrates. Thus, while one volume of serum added to one, two, or three volumes of culture filtrate may still give error due to non-specific group reactions, a proportion of

¹⁴ Myers. *Centralbl. f. Bakt.*, Vol. 28, 1900.

¹⁵ Michaelis. *Deutsche med. Woch.*, 1902.

¹⁶ Norris. *Jour. of Inf. Dis.*, Vol. 1, 1904.

¹⁷ Zupnik. *Zeitschr. f. Hyg.*, 49, 1905.

¹⁸ Kraus. *Wien. klin. Woch.*, 1901, No. 29.

¹⁹ Norris. *Jour. of Inf. Dis.*, Vol. 1, 1904.

one part of serum to 8 or 10 parts of the filtrate will usually eliminate all secondary reactions and prove strictly specific.

An illustration of such an elimination of "partial" or "minor" precipitins by diminution of the amount of the homologous anti-serum is given in the following table taken from the work of Norris²⁰:

ANTICOLI RABBIT SERUM

TABLE III

The precipitating action of the anticoli rabbit serum upon its corresponding filtrates and upon the filtrates of B. N° 1 (hog cholera) and *B. typhosus*.

| Coli filtrate | Anticoli serum | |
|------------------------------|----------------|---|
| 0.5 c. c. | 0.05 | Cloudiness in all tubes in 1 hour at 37.5° C. which increases rapidly. Six hours well-marked precipitation—most copious in tube containing 0.25 serum. Fluid in all tubes becomes clear. |
| 0.5 c. c. | 0.10 | |
| 0.5 c. c. | 0.15 | |
| 0.5 c. c. | 0.25 | |
| B. N°1 filtrate | Anticoli serum | |
| 0.5 c. c. | 0.10 | At 6 hours a slight precipitate in the form of fine granules appears on the sides of the tubes. After 24 hours the precipitate in the tube containing 0.25 c. c. serum compares in amount to that formed in the homologous filtrate with 0.05 c. c. of serum. |
| 0.5 c. c. | 0.25 | |
| B. typh. (Coll) filtrate | Anticoli serum | |
| 0.5 c. c. | 0.10 | Similar reaction obtained to that with B. N° 1 filtrate. |
| 0.5 c. c. | 0.25 | |
| B. typh. (Pfeiffer) filtrate | | |
| 0.5 c. c. | 0.10 | Similar delay in reaction as obtained with B. typh. Coll. |
| 0.5 c. c. | 0.25 | |

And, indeed, though the great practical value of the precipitin reaction has not been in the special field of bacteriology, it has been successfully utilized in the diagnosis of glanders by Wladimiroff,²¹ and constitutes a valuable adjuvant to our methods of determining the biological relationship between micro-organisms.

The production of precipitins against unformed proteins, egg albumen, animal sera, etc., is much more easily accomplished than the production of bacterial precipitins, and three intravenous injections of from 2 to 5 c. c. of the protein at 5 or 6-day intervals usually give rise to a formation of potent precipitins. When a small quantity of the serum of such an animal, taken 9 or 10 days after the third injection, is mixed in a test tube with an equal quantity of a

²⁰ Norris. *Jour. of Inf. Dis.*, Vol. 1, 1904, p. 472.

²¹ Wladimiroff. "Kolle u. Wassermann Handbuch," article on "Glanders," Vol. 5, 2d Ed.

dilution of the protein which was injected, turbidity and rapid flocculation will result. In tests of this kind, unlike the bacterial precipitin tests in which the delicacy of the reaction is ordinarily determined by diminution of the amounts of antiserum, the same object may be more conveniently attained by dilution of the antigen. Thus, in testing the precipitating potency of, let us say, the serum of a rabbit immunized with sheep serum, we would proceed by setting up a series of small tubes, each of which contains a constant amount of antiserum (precipitin), but a progressively diminishing amount of antigen in the same volume—i. e., in dilution with isotonic salt solution. The following example will make this clear:

| Antisheep serum from rabbit | | Sheep serum 0.5 c. c. of following dilutions: | | Precipitation |
|--------------------------------|---|--|---|---------------|
| 0.5 c. c. | + | 1:10 | = | ± |
| 0.5 c. c. | + | 1:100 | = | +++ |
| 0.5 c. c. | + | 1:500 | = | +++ |
| 0.5 c. c. | + | 1:1,000 | = | ++ |
| 0.5 c. c. | + | 1:5,000 | = | + |
| 0.5 c. c. | + | 1:10,000 | = | — |

In this test it will be noticed that the strongest concentration of the antigen (1:10) gave a relatively slight precipitation only. This phenomenon is analogous to the inhibition zones noticed in the case of agglutination and other antibody reactions and will be more especially discussed in a succeeding paragraph.

The delicacy of the above example, moreover, is by no means unusual, and sera have been obtained by careful immunization with which the specific antigen could be detected in dilutions as high as 1 to 100,000 (Uhlenhuth). A serum which will react with antigen dilutions of 1 to 10,000 and 1 to 20,000 is not at all uncommon nor difficult to obtain. Apart from the additional advantage of the specificity of the reaction, therefore, this biological method of detecting proteins is more delicate than that of any of the known chemical methods; neither the Biuret nor Millon's reaction will far exceed a delicacy of 1 to 1,000. By a modification known as the method of Complement or Alexin-fixation, furthermore, the delicacy of the biological reactions can be still further enhanced. This is discussed in detail in another place (see page 212).

The practical value of the precipitin reaction, however, lies, not in the mere detection of protein, but, by virtue of its specificity,²² in the determination of the variety of protein under examination. And

²² Wassermann and Schütze. *Deutsche med. Woch.*, 1900, Vereinsbeilage, p. 178; *Berl. kl. Woch.*, 1901; *Deutsche med. Woch.*, 1902; Bordet, *Ann. Past.*, Vol. 13, 1899; Nolf, *ibid.*, Vol. 14, 1900; Fish, *Medical Courier*, St. Louis, 1900, cited from Wassermann.

here again the specificity, like that of bacterial precipitation, agglutination, and other serum tests, is relative rather than absolute. Thus a serum which has been obtained by the immunization of an animal with human serum may react, not only with human serum, but also with relatively higher concentrations of the sera of some of the higher apes. However, such non-specific partial reactions can be eliminated entirely by employing higher dilutions of antigen. Thus Uhlenhuth,^{23, 24, 25} on the basis of a large experience, has established a standard of antigen dilution at 1 to 1,000, beyond which no "para" or "minor" precipitation will occur. Since potency far exceeding this is easily procured, absolute specificity can be ensured by the very simple precaution of a sufficient dilution.

The most important practical use for the reaction has been found in forensic medicine, where it is possible in this way to determine the species of animal from which have emanated the blood, sperm, etc., found in spots on wearing apparel, weapons, or other articles. The extensive investigations of Nuttall²⁶ upon this subject have incidentally been of much value in furnishing a further method for the determination of zoological species relationships. Nuttall carried out 16,000 precipitin tests, with precipitating sera, upon 900 specimens of blood which he obtained from various sources. He not only confirmed many of the accepted zoological classifications, but shed much light upon a number of disputed points. In working out the tests upon monkeys he found that the reactions carried out with anti-human serum become weaker as the species examined is farther removed from man zoologically. Thus as we read down the column from man to the hapalidæ the precipitate becomes less and less in amount.

Nuttall's Tests with Antihuman Serum. (Nuttall, loc. cit., p. 165.)

ANTHUMAN PRECIPITATING SERUM

| Tested against | Precipitate |
|-------------------------------|--------------------|
| 34 Specimens human blood..... | 100% ²⁷ |
| 8 Simiidæ, 3 species..... | 100% |
| 36 Cercopithecidæ..... | 92% |
| 13 Cebidæ..... | 78% |
| 4 Hapalidæ..... | 50% |
| 2 Lemuridæ..... | 0 |

²³ Uhlenhuth. *Deutsche med. Woch.*, 1900, 1901; *Rob. Koch Festschrift*, 1903.

²⁴ Uhlenhuth and Weidanz. "Kraus u. Levaditi Handbuch," etc., Vol. 2, 1909.

²⁵ Uhlenhuth and Weidanz. *Loc. cit.*, where other publications are summarized.

²⁶ Nuttall. "Blood Immunity and Blood Relationship," Cambridge University Press, 1904.

²⁷ The percentages refer to the volume of precipitate formed on standing for a given time, the amount formed by the antiserum with its specific antigen being taken as 100 per cent. Antigen dilutions correspond throughout.

In another series he finds:

ANTIHUMAN PRECIPITATING SERUM

| Tested against | Precipitate |
|--------------------------|----------------------|
| Man..... | 100% |
| Chimpanzee..... | 130% (loose precip.) |
| Gorilla..... | 64% |
| Ourang..... | 42% |
| Cynocephalus mormon..... | 42% |
| Cynocephalus sphinx..... | 29% |
| Ateles..... | 29% |

Among the primates the highest figures with antihuman serum are given by the chimpanzee. Other bloods than those of the primates gave slight reactions or none whatever with the antihuman serum.

In addition to these results the relationships within the dog family, the horse family, and many other kinships similar to these were confirmed. In every case the precipitin reaction was consistent with the results of other methods of classification, and Nuttall's work is an extremely valuable aid to zoölogists in disputed questions of animal relationships.

These facts are the more surprising in that they demonstrate species differences between the proteins of various animals which are not determinable by known chemical methods. How fundamental these differences are and how delicate the reaction, is further shown by experiments of Uhlenhuth, in which he obtained a specific antihare serum by treating rabbits' with hares' blood, an astonishing result in view of the close zoölogical relations between these animals.

Isoprecipitins, that is, precipitins resulting from the treatment of animals with blood of another individual of the same species, have also been described by Schütze and others. They are not, however, regular in their appearance, nor are they very potent when obtained.

Since the reaction is equally applicable to vegetable proteins, similar investigations on the interrelationship of different varieties of wheat have been carried out by Magnus.²⁸

The methods of performing precipitin tests for forensic or other purposes is extremely simple. Nevertheless, there are a number of theoretical considerations which we must take up in order to make clear the limitations of accuracy and conditions of control which are involved in these reactions. From our discussion of the nature of precipitinogen it follows that blood stains, etc., on linen or articles of any kind will be suitable for precipitin tests even after they have been exposed for considerable periods to unfavorable conditions, that is, an environment in which they are subjected to exposure to light, moderate heat, or drying. Thus blood spots, etc., if kept dry and in

²⁸ Magnus. Cited from Uhlenhuth, *loc. cit.*

the dark, may give positive reactions even after years, as experiments by Uhlenhuth have shown. Meyer²⁹ claims even to have obtained a precipitation with extracts of the material of mummies. One of his specimens was a mummy dating back to the first Egyptian Empire (5,000 years), the other about 2,000 years old. Pieces of the leg and neck muscles of these specimens were chopped up finely, extracted for 24 hours with salt solution, then filtered until clear. With antihuman serum they gave turbidity after one hour at 37.5° C.

Under conditions of putrefaction, of course, the precipitinogen is more rapidly destroyed, though blood putrefies with surprising slowness, even if, as in our own experiments, the conditions of moisture, temperature, and reinoculation with putrefactive bacteria are constantly observed. Under such conditions a weak reaction may be obtained after as long as a month or six weeks.

In carrying out the tests with any material it is first necessary to get it into clear solution, a result which is best accomplished by soaking it in a small quantity of isotonic salt solution. Preliminary to this it is always necessary to scrape off a bit of the specimen and examine it microscopically to discover, if possible, whether blood cells, sperm, or other cellular constituents can be detected. The infusion in salt solution should be continued for several hours—if necessary for 12 to 24 hours. After the first few hours in the incubator the material should be placed at room or refrigerator temperature so that the yield in unchanged protein may not be diminished by the action of bacterial growth. After extraction the solution may be filtered in order to clear it, but often mere centrifugation suffices for this purpose. The concentration of antigen in such an extract is always an uncertainty, but may be determined with sufficient accuracy for practical purposes by shaking and observing the formation of a lasting foam. Protein solutions will show foam on shaking in dilutions as high as 1 to 1,000, and if the original amount of salt solution used in washing out the material is properly gauged to the amount of blood available in the stain, and the solution shaken and observed for the formation of foam, it is usually a simple matter to obtain a final concentration approximating one to one thousand.³⁰

The antiserum which is used should be of such a potency that preliminary titration with the specific antigen, diluted 1 to 1,000, should give an almost immediate cloudiness at room temperature.

By testing this serum against a number of other varieties of

²⁹ Meyer. *Münch. med. Woch.*, Vol. 51, No. 15, 1904.

³⁰ If there is enough material, the amount of dissolved protein can be also approximately gauged by adding to a little of it a drop of acid, boiling and observing the heaviness of the cloud which forms. A control test of a known dilution of the suspected variety of blood can be made at the same time and the heaviness of this cloud compared with that in the test solution.

protein—dog serum, beef serum, etc.—it must be determined that the precipitin in this case is strictly specific.

The reaction can be observed with greater delicacy if it is first set up by the method recommended by Fornet and Müller,³¹ which we may speak of as the “ring test.” The antiserum is put into the tubes and the solution to be tested is allowed to flow slowly over this—as in Heller’s nitric acid albumin test. At the line of contact between the two a fine white ring will rapidly appear, thickening and growing heavier as the preparation is allowed to stand. After taking the final readings from such a test, let us say after an hour or so, it is well to shake up the tubes, set them away in the ice-chest, and again read the amount of precipitates formed in the various tubes the next morning. Since every test of this kind necessitates a number of controls, the following example will serve as a basis for discussion:

Forensic Blood Examination

Material: Blood spot on trouser pocket, washed up in salt solution. Clear after paper filtration.

Antiserum: Rabbit treated with three intravenous injections, 2, 5, and 5 c. c. of human serum at six-day intervals; bled on tenth day after last injection. This serum has been titrated against human serum and gives precipitation in dilutions up to one to ten thousand. With one to one thousand there is clouding which begins in three minutes and is very distinct in eight minutes, at room temperature.³²

Test

| | | |
|---|--|-----------|
| Tube 1. Known human serum 1 to 1,000 . . . | 1.0 c. c. + Antiserum . . . | 0.2 c. c. |
| Tube 2. Unknown solution to be tested . . . | 1.0 c. c. + Antiserum . . . | 0.2 c. c. |
| Tube 3. Unknown solution to be tested . . . | 1.0 c. c. + Normal rabbit serum | 0.2 c. c. |
| Tube 4. Salt solution | 1.0 c. c. + Antiserum . . . | 0.2 c. c. |
| Tube 5. Unknown solution | 1.0 c. c. + Salt solution . . . | 0.2 c. c. |

In this test, if the original material was human blood, tubes 1 and 2 should show ring formation within 5 minutes—while the other tubes remain clear. In addition to these controls it is well to be sure that the test extract is neither strongly acid nor alkaline, and that, as Uhlenhuth suggests, the material from which it is extracted does not contain other substances which can give precipitates by themselves when added to serum. This is especially necessary in the case of cloth fabrics, and a recent instance in our own experience has suggested to us the possibility that such materials may also contain colloidal dye stuffs or other extractable substances which can cause inhibition of the precipitation. In an apparently positive case the reactions with a blood extract from trouser cloth were sufficiently heavy, but regularly delayed, as in the flocculation of such

³¹ Fornet and Müller. *Zeitschr. f. Hyg.*, Vol. 66, 1910.

³² A mixture of too specific antisera should never be used, since such sera may often precipitate each other for reasons that are discussed below.

colloidal suspensions as arsenic trisulphide in the presence of a protective colloid.

In the ordinary criminal or civil case which would come under consideration for precipitin tests the spots or stains are made by blood as it flows from the wound and unchanged by chemical or physical agencies except as these are encountered afterward, by exposure. In the case of meat inspection, in which the precipitin test is useful in detecting admixtures of horse flesh, dog flesh, or other less desirable varieties of meat, in sausages, chopped meat, etc., it often happens that such procedures as heating or smoking may vitiate the results of precipitin reactions. It is of practical importance, therefore, that we should know exactly what the effects of heating (boiling) may be upon precipitinogen. Moreover, this question possesses considerable theoretical interest since the coagulation of proteins by heat seems to involve chiefly a physical rather than a chemical change.

Cohnheim³³ says in discussing this question: "It is still unclear what the changes are that take place in coagulation. It may be that there is merely an intramolecular 'Umlagerung'—or there may be cleavage; or the process may be comparable to the flocculation of colloidal clay emulsions by salts. . . . With coagulation all proteins have lost the differences which they possess in the native state in respect to solubility or precipitability by salts. Physically all coagulated proteins are alike; they are no longer native proteins, and without further decomposition are insoluble. Chemical differences, however, variations of composition, and the cleavage products which they yield still distinguish them."

The question has been experimentally approached by Obermeyer and Pick³⁴ in connection with their general investigations upon the influence of chemical and physical alterations upon precipitinogen. They found that precipitin produced with unchanged (native) beef serum does not react with heated beef serum, even if immunization was prolonged and a very potent serum was produced. On the other hand, when animals were immunized with beef serum which had been boiled for a short time ("Kurz aufgekocht"³⁵) the precipitin so produced reacted, not only with native beef serum, but also precipitated the boiled serum and a whole row of split products which give no reaction to normal precipitin. The "coctoprecipitin" so produced, furthermore, was found by them to be specific, acting only upon beef protein or its derivatives.

³³ Otto Cohnheim. "Chemie der Eiweiss Körper Vieweg Braunschweig," 1900, p. 8.

³⁴ Obermeyer and Pick. *Wien. kl. Woch.*, 12, 1906.

³⁵ Sera or other proteins used in such tests are boiled in dilutions of 1 to 10 or more, in order to avoid the formation of heavy flakes which cannot be injected. Boiled in sufficient dilution, an opalescent suspension is formed which easily passes through the syringe.

It is immediately evident that these investigations are closely analogous to those of Joos and others on the agglutinins. The antiserum produced with the heated antigen here again reacts both with the native and with the heated antigen, whereas the antiserum produced with the native unheated antigen reacts only with the unheated. The "heat-precipitins" therefore may be also called "umfänglicher"—the term applied by Paltauf to the agglutinins produced with heated bacteria.

Schmidt,³⁶ who has studied the problem extensively, finds that heating serum protein to 70° C. for as long as 30 to 60 minutes alters its precipitability by "native precipitin" (precipitin produced by immunization with native unheated serum) only in so far as it diminishes the delicacy of the reaction by 10 to 30 per cent., and that heating to 90° C. for as long as an hour does not render it entirely non-precipitable, so that protein so treated may yet be detectable by ordinary specific precipitins produced by injections of unheated serum, though the delicacy of the reaction is lessened. Boiling, according to Schmidt, renders the antigen no longer precipitable by such "native precipitin," but, on the other hand, it does not seem to destroy its antigenic property of inciting precipitins on injection into animals. Fornet and Müller, on the other hand, claim that even boiled protein can be detected by "native precipitins," though the reaction is only about one-tenth as delicate as it is with unheated protein.

Schmidt studied these relations especially as they affect the performance of specific precipitin reactions in the identification of boiled meat. He found that when he immunized rabbits with serum protein that had been heated at 70° C. for 30 minutes the antiserum so obtained gave strong and practically useful reactions with its specific antigen even if this had been boiled. Since "native precipitin" gives weak reactions only with such a boiled protein, Schmidt recommends the use of the "70° precipitin" (produced by injections of heated serum) for tests in which a heated antigen is to be identified.

He states, however, that very prolonged heating may so completely coagulate the antigen that none of it can be gotten into "solution" (suspension), and in such cases results can be obtained neither with the "native" nor with the "70° precipitin." He has attempted, therefore, to find a method whereby even such entirely insoluble proteins may be identified, and claims to have succeeded by preparing what he calls his "heat-alkali-precipitin."³⁷ He di-

³⁶ Schmidt. *Biochem. Zeitschr.*, 14, 1908; also *Zeitschr. f. Imm.*, Vol. 13, 1912.

³⁷ "Native precipitin" = precipitin produced by injections of normal unheated serum.

"70° precipitin" = precipitin produced by injections of serum heated to 70° C. for 30 minutes.

lutes serum with equal parts of isotonic salt solution and heats it to 70° C. for 30 minutes in a water bath. To 60 c. c. of such a solution he now adds 10 c. c. of $\frac{n}{1}$ NaOH, and continues heating for 15 to 20 minutes. At the end of this time he neutralizes with HCl, cools, and injects 20 c. c. intraperitoneally into rabbits. (The neutralization is not absolutely necessary.) Five or more injections yield a serum sufficiently potent for use.

A precipitin so produced will, according to Schmidt, react specifically with heated proteins, and also with protein which has been solidly coagulated and brought into solution by means of NaOH and heat. It will not, however, react with normal unheated antigen.

He tested this by coagulating horse serum by boiling for 3 hours. The coagulum was washed with salt solution, dried, and powdered. Tests were then made to prove that this powder was entirely insoluble in NaCl solution. A little of it was then treated with 10 c. c. of salt solution containing enough NaOH to correspond to an $\frac{n}{1}$ solution. The exposure was continued for 20 minutes in a water bath at 60° to 70° C. Before the entire mass was dissolved the solution was filtered and neutralized with $\frac{n}{20}$ HCl.

The rather complicated relations described by Schmidt are easily surveyed in the following protocol taken from his work:

TABLE I

(W. A. Schmidt, *Zeitschr. f. Imm.*, Vol. 13, 1912, p. 173)

| Solution of | Native precipitin | Heat (70°) precipitin | Heat-alkali-precipitin |
|--|-------------------|-----------------------|---------------------------|
| Native serum..... | Strong reaction | Good reaction | 0 (very slight turbidity) |
| 70° serum (heated 30 min.) | Good reaction | Strong reaction | Strong reaction |
| 100° serum (heated 30 min.)..... | 0 | Good reaction | Strong reaction |
| 70° serum treated with NaOH (used to produce heat-alkali-precipitin) . | 0 | 0 | Strong reaction |
| Boiled insoluble serum, brought into solution with NaOH..... | 0 | 0 | Good reaction |
| Native serum treated with NaOH in the cold..... | 0 | 0 | Good reaction |

Schmidt speaks of the "heat-alkali-precipitin" also as "alkali-albuminate-precipitin." It can be produced only if the NaOH treatment of the serum is cautiously performed. If the sodium hydroxid is allowed to act too vigorously in strong concentrations or for too long a time the antigen is completely destroyed, is no longer pre-

ipitable, and no longer produces precipitin when injected into animals.

The striking feature of these experiments is that they show a gradual alteration of the protein first by heat, then by alkali and heat, in such a way that the antigenic properties are changed but not destroyed. Each precipitin, moreover, seems to react most strongly with the particular antigen-alteration which produced it, and, according to Schmidt, retains its species specificity. This is not the case with the iodized proteins and nitroproteins and diazoproteins produced by Obermeyer and Pick.³⁸ Here iodized beef protein injected into animals produced a precipitin which reacted with the iodized protein, not only of the beef, but also similarly altered proteins of other animals—and the same was true of the nitro and diazo modifications.

Although the experiments of Schmidt have great theoretical value, their practical utilization must depend upon the degree of specificity possessed by the heat-precipitins or the heat-alkali-precipitins. In Obermeyer and Pick's original investigations we have seen that they found the precipitin produced with heated serum as strictly specific as that induced by native serum. This has also been the experience of Schmidt. Fornet and Müller,³⁹ on the other hand, report that the precipitins produced by them with heated muscle-protein were not as strictly specific as those produced with the unheated—in that the former gave precipitates, not only with homologous protein solutions, but with foreign proteins in moderate concentration as well. In experiments carried out by the writer with Ostenberg⁴⁰ it was attempted to determine whether or not precipitins could be produced by injecting animals with protein that had been boiled, and if so what the action of these substances would be upon boiled proteins. Contrary to the results of Fornet and Müller, it was actually found that sera boiled for 3 to 5 minutes injected into rabbits induced precipitins which acted upon boiled proteins, but at the same time it was determined that the antibodies so produced were no longer strictly specific. The protocol given at the top of the next page will illustrate these experiments.

Summarizing these results together with those of Fornet and Müller and of Schmidt it would seem that the injection of boiled proteins induces precipitins which no longer act on native antigen, which act powerfully on boiled antigen, but are no longer strictly specific. This seems to us of great theoretical interest as showing an alteration by heating in the species adherence of the antigen. Practically, therefore, precipitins produced with boiled protein are of little value, and forensic determinations of boiled proteins should

³⁸ Obermeyer and Pick. *Wien. klin. Woch.*, No. 12, 1906.

³⁹ Fornet and Müller. *Zeitschr. f. Hyg.*, Vol. 66, 1910.

⁴⁰ Zinsser and Ostenberg. *Proc. N. Y. Pathol. Soc.*, 1914.

Experiments on Cocto-precipitin. Table II (March 23, 1913).

Cross titrations—dilutions of sera in salt solution boiled 5 minutes, precipitated with antisera produced by injections with similarly boiled material.

The readings here indicated were taken by "ring" test at the end of 30 minutes.

| Dilution | Beef serum vs. anti-beef precipitin | Beef serum vs. anti-dog precipitin | Beef serum vs. anti-sheep precipitin | Dog serum vs. anti-dog precipitin | Dog serum vs. anti-beef precipitin | Dog serum vs. anti-sheep precipitin | Sheep serum vs. anti-sheep precipitin | Sheep serum vs. anti-dog precipitin | Sheep serum vs. anti-beef precipitin |
|---------------------------------|-------------------------------------|------------------------------------|--------------------------------------|-----------------------------------|------------------------------------|-------------------------------------|---------------------------------------|-------------------------------------|--------------------------------------|
| 1:20 | + | + | + | ++ | — | + | ++ | ++ | + |
| 1:50 | +++ | + | +++ | ++ | — | ++ | +++ | + | +++ |
| 1:100 | +++ | + | +++ | ++ | — | + | ++ | + | ++ |
| 1:500 | ++ | — | + | + | — | + | + | — | — |
| 1:1,000 | ± | — | — | ± | — | ± | ± | — | — |
| Controls of boiled serum alone* | | | | | | | | | |
| 1:20 | — | | | — | | — | | | |
| 1:50 | — | | | — | | — | | | |
| Serum control | | | | | | | | | |

* These controls were necessitated by the fact that the boiled serum suspensions were themselves turbid and occasionally showed slight settling on standing.

be done, as advised by Schmidt, by the "70° precipitins," or with native precipitin as practiced by Fornet and Müller.

The specificity which is the basis of the practical value of the reactions that we have discussed is spoken of as "species" specificity since it has been found that the blood serum of rabbits or other animals into which the serum of another animal has been injected reacts, not only with the homologous blood serum, but also with extracts of the various organs of the particular species of animal which furnished the serum. Thus if we immunize rabbit, let us say, with sheep serum the resulting precipitin will react, not only with sheep serum, but also with extracts of sheep spleen, sheep liver, etc. It seems that every species of animal possesses throughout its tissues a particular variety of protein, fundamental to its general metabolism and peculiar to its species. On the other hand, we have seen in the preceding discussions how chemically slight the changes in a protein may be which can alter materially its antigenic nature, and it is a logical deduction that different organs of the same animal might contain antigenic constituents qualitatively different from the general serum protein. There are undoubtedly in many organs protein complexes which are peculiar to them and not present in other organs, and it would be reasonable to expect therefore that immunization with separate organ substances would lead to the production of sera of specific precipitating power for the protein of that particular kind of organ. This is not ordinarily obtainable, how-

ever, because it has been impossible to isolate from organs their peculiar, characteristic proteins, and immunization of animals with organ extracts or solutions has necessarily implied the injection of much blood protein and other albuminous material of a character general to many organs of the animal, i. e., to the species. These quantitatively overshadow the organ-specific substances which may be present, and give rise, therefore, to a "species" precipitin. That "organ specificity," however, is a fact has been shown by the experiments of Uhlenhuth with the protein of the crystalline lens of the eye. Immunization with this substance induces a precipitin which does not react with the serum of the animal from which the lens was taken, but *does* react, not only with the crystalline lens proteins of this species of animal, but also with crystalline lens proteins in general, though taken from another animal species. Analogous to this are the experiments of von Dungern and others upon the protein derived from the testicle.

In both of these cases, as well as in other less sharply defined examples, the specificity is attached, not to the species of animal, but rather to the nature of the organ from which the particular protein is derived. These facts—first ascertained by means of the precipitin reaction—have been recently confirmed by means of the reaction of anaphylaxis by Uhlenhuth and Haendel, and by Kraus, Doerr, and Sohma. (See chapter on Anaphylaxis.) They have been discussed, moreover, in connection with the problem of specificity in general.

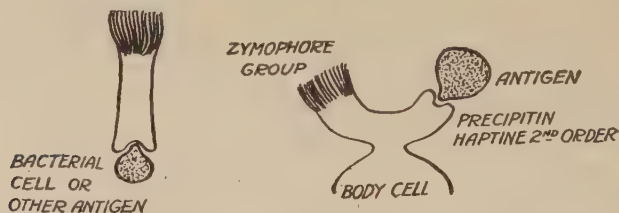
Biologically they probably signify that, although there are fundamental species differences between the general body proteins of various animals, there are still, in certain highly specialized organs, varieties of protein which, possibly because of functional exigencies, have developed similar chemical characteristics. These have been determinable by our present methods, however, only for organs like the lens, the testicle, and the placenta from which the organ-specific protein can be gotten in a relatively pure state. The pathological importance of these phenomena lies in the fact that, although guinea pig serum injected into a guinea pig will not give rise to antibodies, lens protein apparently will do so—an observation which opens the possibility of autocytotoxins. The significance of this is indicated in such investigations as those of Römer,⁴¹ who, using the complement-fixation technique to determine antibody, found that the serum of adult human beings possessed antibodies for their own lens protein, but that such antibodies were absent in the sera of children.

The study of agglutination and that of precipitation reveal, throughout, a close similarity between the two reactions, and indeed in physical principles they are probably the same, although the one

⁴¹ Römer. *Klin. Monatsbl. f. Augenheilkunde*, Sept., 1906. Ref. from "Weichhardt's Jahresber.," Vol. 2, 1906, p. 348.

(agglutination) consists in the flocculation of large particles in suspension—the bacteria—while in the other the precipitation is one of smaller units—the precipitable colloidal particles of the protein solutions. This phase of the subject will be more thoroughly discussed directly.

Meanwhile, it is noticeable also that, even without drawing the physical parallel between the two reactions, there is much in the behavior of the antibodies—the agglutinins and the precipitins as conceived by Ehrlich, which led him and his school to attribute to them a similar receptor structure. Like the agglutinins, the precipitins are not inactivated by 56°C ., but when once rendered ineffectual by higher temperatures (70°C . or over) they can no longer be reactivated by the addition of fresh normal serum. For this reason chiefly Ehrlich has conceived that both agglutinins and precipitins are “haptines” of the second order.



SCHEMATIC REPRESENTATION OF EHRlich's VIEWS ON THE STRUCTURE OF PRECIPITINS.

Ehrlich assumes that when dissolved protein substances—ordinarily suitable for body nutrition—are injected into animals, they become anchored to the cells by such receptors of the second order. When overproduction occurs in response to repeated stimulation of the cells by consecutive injections (see Side-Chain Theory), these haptines of the second order circulate as agglutinins or precipitins. Since they act without the apparent coöperation of alexin, he supposes that they carry within themselves the “zymophore,” or ferment groups, by means of which the agglutination or coagulation is accomplished. It is this zymophore group which, it is assumed, accomplishes the digestion of the foreign protein before its assimilation, when these receptors are still parts of the living cell.

Thus the conception of precipitins is identical with that formulated by the same school concerning the agglutinins, and the deductions from these premises have been essentially similar. Thus, analogous to the conditions prevailing in agglutination, Pick,⁴² and Kraus and v. Pirquet⁴³ have shown that when precipitating serum is inactivated by heat, and then is added to bacterial filtrates, it will

⁴² Pick. “Hofmeister's Beiträge,” Vol. 1, 1902.

⁴³ Kraus and von Pirquet. *Centralbl. f. Bakt.*, Vol. 32, 1902.

prevent their subsequent precipitation by active precipitin. An illustration of this is found in the following protocol taken from the paper by Kraus and v. Pirquet (*loc. cit.*, p. 69).

- (a) 5 c. c. cholera filtrate + 0.5 c. c. inactiv. (60°) cholera serum = no precipitate after 10 hours at 37° C.
 After 10 hours add 0.5 c. c. active cholera serum = no precipitate.
 (b) Omitted.
 (c) Omitted.
 (d) 5 c. c. cholera filtrate + 0.5 c. c. active cholera serum = after 10 hrs. typical precipitate.

From this it was concluded that heat may destroy the zymophore or coagulating group of precipitins, leading to the formation of "precipitinoids" which, like agglutinoids, may have a higher affinity for the antigen than is possessed by the uninjured antibody.

Subsequently there were opposed to these views the physical interpretations which have been outlined sufficiently in the section on Agglutination (see p. 240). In the case of precipitation the analogy between colloidal reactions and the serum phenomena is fully as striking as in the former, an analogy in the delineation of which the first credit belongs to Landsteiner,^{44 45} and important further contributions have been made by Neisser and Friedemann, Porges, Gengou, and a number of others. As in agglutination and colloidal flocculation, the presence of salts (electrolytes) fundamentally influences the occurrence of precipitin reactions; and in both colloidal and precipitin reactions the relative concentration of the reacting bodies is paramount in determining whether or not precipitation takes place. In this connection the most frequently observed inhibition occurring in serum precipitations is that which is caused by an excess of antigen. An example of this is as follows:

| Sheep serum 0.5 c. c. | | Antisheep rabbit serum | Precipitate |
|-----------------------|---|------------------------|-------------|
| 1:10 | + | 0.5 c. c. | — |
| 1:50 | + | 0.5 c. c. | ± |
| 1:100 | + | 0.5 c. c. | ++ |
| 1:500 | + | 0.5 c. c. | +++ |
| 1:1,000 | + | 0.5 c. c. | ++ |
| 1:5,000 | + | 0.5 c. c. | + |

This is entirely analogous to the inhibition which may occur when, let us say, a weak gelatin solution is added to a colloidal suspension of arsenic trisulphid; or blood serum is added to mastic or arsenic suspensions. In both cases inhibition zones appear which

⁴⁴ Landsteiner and Jagie. *Münch. med. Woch.*, No. 18, 1903; No. 27, 1904; *Wien. kl. Woch.*, No. 3, 1904.

⁴⁵ Landsteiner and Stankovic. *Centralbl. f. Bakt.*, Vols. 41 and 42, 1906.

show that the relative quantities of the two reacting bodies are quite as significant as their chemical or physical constitution in determining the occurrence of flocculation. This, according to Bechold, Billitzer,⁴⁶ and others depends upon the fact that the reason for flocculation is one of electrical charge. One hydrosol—say arsenic trisulphid—can be flocculated by the oppositely charged colloidal aluminium hydroxid, but this will occur only when the quantitative relations are properly adjusted. If one or the other is in excess, no flocculation may occur, and, if subjected to a direct current, both colloids, though ordinarily wandering in opposite directions, will now wander in that of the one which is now present in the largest amount. We will not elaborate here upon the causes for this, since they have been indicated in the section on Agglutinins, and are set forth more accurately by Prof. Young in the special chapter on Colloids.

This effect of quantitative proportions would explain not only the absence of precipitation in the presence of too much antigen, but also the converse phenomenon, already mentioned, that precipitation may be inhibited when the precipitin is in excess.

The fact that heated precipitating serum when added to its antigen not only does not cause flocculation, but may even prevent subsequent precipitation by active precipitin, also finds its analogy in colloidal reactions in the so-called protective colloids. Thus arsenic trisulphid may be protected from precipitation by gelatin, if a small amount of gum arabic is added, and the analogy has been brought even closer by Porges,⁴⁷ who showed that heated serum will protect mastic suspension from precipitation by normal serum. This observation of Porges is so closely similar to the results obtained by Kraus and v. Pirquet and others on the inhibition of precipitation by heated precipitating serum that it would seem, on first consideration, effectually to refute the conception of "precipitoids."

However, it does not explain the specificity of such inhibition on the part of heated precipitating serum, as reported by Kraus and v. Pirquet, an observation which is one of the strongest arguments in favor of the derivation of the inhibiting factor from the specific precipitin (a precipitoid).⁴⁸

In spite of the strong evidence in favor of the colloidal interpretations, such contrary evidence, brought forward by careful and

⁴⁶ Billitzer. Cited from Bechold, "Die Kolloide, etc.," p. 79.

⁴⁷ Porges. Chapter on "Colloids and Lipoids" in "Kraus u. Levaditi Handbuch," Vol. 1.

⁴⁸ Although normal sera may gradually precipitate on standing, this takes place much more rapidly in precipitin-sera. The spontaneous precipitation of normal sera as well as of those under consideration is analogous to what Bechold and others call the "ageing" (altern) of colloidal suspensions, which, though originally stable, will eventually settle out, even in the presence of protective colloids.

experienced workers, must be borne in mind and positive acceptance of the colloidal explanations, however attractive, must be withheld until much further investigation has been done.

Another important and interesting phase of the study of precipitins is that associated with the occasional presence in the same serum of remnants of antigen and of precipitins which, though present side by side, do not unite to form precipitates. This condition is frequently seen in such sera as those produced by Fornet and Müller⁴⁹ for rapid precipitin production for forensic work, a method in which the foreign serum is injected into rabbits in large amounts (2 to 10 c. c.), on consecutive days, and the animals are bled 6 to 8 days after the last injection. That such sera contain both antigen and antibody is shown by the fact that, though clear when taken, they will show precipitation not only when mixed with dilutions of the antigen, but also when added to homologous precipitating sera.⁵⁰

This phenomenon has been noticed by Linossier and Lemoine,⁵¹ Eisenberg,⁵² Ascoli,⁵³ and others, and has been extensively studied by von Dungern.⁵⁴ Gay and Rusk⁵⁵ have recently observed it in connection with the rapid method of precipitin production of Fornet and Müller, and have noted that such sera, although containing both antigen and precipitin, do not possess complement-fixing properties. According to Uhlenhuth and Weidanz,⁵⁶ the antigen may persist in the sera of protein-immunized animals, in demonstrable amounts, as long as fifteen days after the last injection, and it is constantly present during this period, but in progressively diminishing amounts.

We are thus confronted by the apparently paradoxical phenomenon of the presence in these sera, side by side, of an antigen and its homologous precipitin, incapable of reacting with each other, although each of them readily reacts with precipitin or antigen, respectively, when these are added from another source.

Many attempts have been made to account for this. A number of observers, notably Eisenberg, have concluded from extensive an-

⁴⁹ Fornet and Müller. *Zeitschr. f. biol. Technik u. Methodik*, Vol. 1, 1908.

⁵⁰ For instance, a rabbit was injected on three consecutive days with sheep serum. It was bled on the fifth day after the last injection. The serum was clear when taken, but a precipitate was formed when it was added to sheep serum and also when it was added to serum from another rabbit similarly treated and containing sheep serum precipitin.

⁵¹ Linossier and Lemoine. *C. R. de la Soc. de Biol.*, 54, 1902.

⁵² Eisenberg. *Centralbl. f. Bakt.*, 34, 1903.

⁵³ Ascoli. *Münch. med. Woch.*, Vol. 49, No. 34, 1902.

⁵⁴ Von Dungern. *Centralbl. f. Bakt.*, 34, 1903.

⁵⁵ Gay and Rusk. "Univ. of Cal. Public. in Pathology," Vol. 2, 1912.

⁵⁶ Uhlenhuth and Weidanz. "Praktische Anleitung zur Ausführung, etc.," Jena, 1909.

alyses of quantitative relationships, both of agglutinin and precipitin reactions, that these take place according to the laws of mass action. In consequence, in addition to the combined precipitin-antigen complex present in all mixtures of the two, there should also be present free dissociated fractions of each, in amounts dependent upon relative concentrations. This might explain conditions such as those described above.

Von Dungern, whose paper forms one of the most extensive studies of the phenomenon with which we are concerned, does not believe that precipitin reactions can follow the laws of mass action, and explains the simultaneous presence of precipitin and antigen in the same serum by assuming a multiplicity of precipitins. He believes that every proteid antigen contains a number of related partial antigens which give rise in the immunized animal each to a partial precipitin. In sera in which both antigen and precipitin are found side by side and free, he believes that the antigen is of a nature that has no affinity for the particular partial precipitin present with it. He says: "Auch hier handelt es sich nicht um zwei reaktionsfähige Körper, deren Verbindung aus irgend welchen Gründen unterbleibt, sondern um Substanzen, welche keine Affinität zu einander besitzen. Die betreffenden Kaninchen haben zu dieser Zeit noch nicht alle möglichen Teilpräzipitine gebildet, sondern nur einzelne derselben. Diese zunächst produzierten, nur auf bestimmte Gruppen der präzipitablen Eiweisskörper passenden Partialpräzipitine sind es, welche nach der Absättigung aller zur Verfügung stehenden zugehörigen Gruppen der präzipitablen Substanz in Serum nachweisbar werden. Daneben bleibt aber ein anderer Teil der präzipitablen Substanz, der keine Affinität zu dem gebildeten Präzipitin besitzt, bestehen, solange bis ein anderes Partialpräzipitin von den Kaninchenzellen geliefert wird, welches sich mit Gruppen der in Lösung geliebten Eiweisskörper vereinigen kann."

Zinsser and Young⁵⁷ have also studied these phenomena and have attempted to explain them on the basis of protective colloidal action. In considering the theories that have been advanced to explain these occurrences, the conception of mass action as accounting for the simultaneous presence of the two reacting bodies in the same serum seemed entirely incompatible with our own observations and with those of Gay and Rusk, that these sera do not of themselves fix alexin. Were the conception of the manner of union of these two reagents, according to the laws of mass action, representative of the true state of affairs, it would be necessary to assume the presence, in such sera, not only of the two reacting bodies free and dissociated, but also of a definite quantity of the united complex of the two, a state of equilibrium being established. If this were the case the sera should, in agreement with all experience on the phenomenon

⁵⁷ Zinsser and Young. *Jour. of Exp. Med.*, 1913, Vol. 17.

of complement fixation, exert definite complement-binding power. Moreover, it has not been experimentally shown that colloidal substances react in accordance with the laws of mass action as observed for simpler chemical substances.

As regards the opinion of von Dungern, this seemed incompatible with another occurrence, observed by many writers, namely, that such sera, although clear at first, eventually, after prolonged standing, *do* actually precipitate spontaneously; that is, the union of the precipitin and the precipitinogen *does* actually take place, but goes on with extreme slowness.

Now a notable and strange feature of this phenomenon is the fact that two such sera, both containing antigen and precipitin, but neither of them precipitating by itself, will precipitate each other when mixed. For this reason Uhlenhuth has advised against the use of mixtures of precipitin sera for forensic tests. For it is not unusual that precipitin sera, even when produced by the slow method, may contain traces of antigen, and this may lead to precipitate formation if such a serum is mixed with another homologous precipitin and thereby simulate a positive forensic test.

In seeking analogy for this serum phenomenon with the various colloidal suspensions, the problem consisted in protecting two mutually precipitating colloids by a third, and this in such proportions that the mixing of two such protected suspensions, each containing all three of the elements, would be followed by precipitation. This was obtained by the use of gum arabic, gelatin, and arsenic trisulphid. Thin emulsions of gelatin will precipitate arsenic trisulphid suspensions. Small amounts of gum arabic will act as a protective agent, preventing the precipitations.

The amount of the protecting substance necessary to prevent precipitation in any one mixture varies apparently with every change in the relative proportions of the two. Thus a considerable number of mixtures of the three can be made which will remain stable for days, the actual and relative quantities of the three ingredients differing in each of the mixtures. When two such mixtures are poured together, in many cases precipitation will result, varying in speed and completeness, according to the particular quantitative relationship arrived at in the mixture.

An example of such an experiment follows:

Two solutions of colloidal arsenic sulphid were prepared, one containing 1 gm. per liter, the other containing 5 gm. per liter. With Kahlbaum's "Gold-ruck" gelatin a solution containing 1 gm. per liter was prepared. A solution of gum arabic was prepared which contained 10 gm. per liter, this being made stronger than the gelatin solution to avoid too great dilution in the final mixtures. The gelatin solution was prepared twenty-four hours before being used, as freshly prepared gelatin has but slight precipitating power for arsenic sulphid, this power appearing to increase greatly with the ageing of the solution.

For the purpose of demonstrating this analogy two protected solutions were prepared as follows:

Solution 1.—This consisted of 2 drops of gum arabic, 2 c. c. of gelatin, and 5 c. c. of the weaker arsenic solution.

Solution 2.—This consisted of 10 drops of gum arabic, 1 c. c. of gelatin, and about 4 c. c. of the stronger arsenic solution.

In each case the arsenic sulphid was added until there were signs of increasing opalescence or turbidity, this being done in order that the two solutions should each be as little overprotected as possible.

Portions of the two solutions were then mixed in equal proportions. In the course of a few minutes the mixture was noticeably more turbid than either of the original solutions. This turbidity continued to increase quite rapidly, and on the following morning after about sixteen hours of standing, the mixture was found to be completely flocculated out, while the original protected mixtures remained unprecipitated and showed about the same degree of opalescence as on the preceding night. The same condition of affairs was found to have persisted after five days. On the fifth day the less concentrated of the clear protected suspension began to settle out, and was completely precipitated within twenty-four hours. The other remained clear for four days more, but on the ninth day it began to precipitate slightly, the precipitation remaining incomplete.

In these cases it appears, therefore, that a complete analogy to the observed conditions of the serum reactions has been found, and that all data observed in connection with sera in which antigen and precipitin are found side by side without reacting can be most simply explained on the conception of protective colloid action. Moreover, the chemical nature of the substances involved seems to add weight to this point of view.

These relations have been gone into here at some length, since they seem to us to possess considerable theoretical and practical significance. For it may be that the presence of a protective colloid may, by inhibiting the union of antigen and precipitin within the body, protect the animal from intoxication during the early stages of immunization when antigen and antibody are present simultaneously for longer or shorter periods. Were union between the two possible at such times in the circulation, an assumption necessitated both by the hypotheses of mass action and of multiplicity of precipitins, there would probably be an absorption of complement by these complexes, with, as shown by Friedberger, a consequent formation of powerful toxic products. (See chapter on Anaphylaxis.) It is not impossible by any means, therefore, that the injection of antigen in an animal in which such a balance has been established may

lead to a sudden elimination of the colloidal protective action, union of the antigen and antibody, and, by the mechanism just outlined, anaphylactic shock.

The fact, moreover, that mere heating will change the precipitating action, which certain sera have on inorganic colloids, to a protective one seems to show that this latter function may justly be associated with delicate physical or chemical alterations of animal sera.

Furthermore, this point of view is strengthened by the fact that the mutual precipitation of sera, such as those described, takes place slowly, as does the mutual precipitation of two protected colloidal mixtures, in contradistinction to the more rapid precipitation which takes place when any of these sera is added to an antigen dilution, where the element of protection may be assumed to be practically eliminated by more extensively changed quantitative relations.

CHAPTER XI

PHAGOCYTOSIS

EARLY investigations into the fate of bacteria within the infected animal body were largely carried out by pathological anatomists, and the observation of the presence of micro-organisms within the cells of the animal and human tissues was definitely made as early as 1870. Hayem,¹ Klebs,² Waldeyer,³ and others, saw leukocytes containing bacteria but failed to interpret this in the sense of possible protection. The process was regarded rather as a means of transportation of the bacteria through the infected body, or it was assumed that possibly the micro-organisms had entered these cells because of the favorable nutritive environment thus furnished.

The first to suggest that such cell ingestion might represent a method of defence was Panum,⁴ who referred to it as a vague possibility. A similar but more convinced expression of this opinion was made in 1881, according to Metchnikoff,⁵ by Roser in explaining the resistance of certain lower animals and plants against bacteria. But Roser brought no experimental support for his contention, and little attention was paid to his assertion.

The significance of cell ingestion as a mode of protection against bacterial invasion, therefore, was hardly more than a vague suggestion when Metchnikoff, who, though a zoölogist, had become intensely interested in the problem of inflammation, began to experiment upon the cell reaction which followed the introduction of foreign material, living or dead, into the larvæ of certain starfishes (*Bipinnaria*).

Pathologists, at this time, held complicated views of inflammation which involved complex coördinated reactions of vascular and nervous systems, and Metchnikoff's primary purpose was to observe reactions to irritation in simple forms devoid of specialized vascular or nervous apparatus. He noted in these transparent, simple forms of life that the foreign particles were rapidly surrounded by masses of ameboid cells and reached a conclusion which, in his own words, is expressed as follows:

¹ Hayem. *C. R. de la Soc. Biol.*, 1870.

² Klebs. *Pathol. Anat. der Schusswunden*, 1872.

³ Waldeyer. *Arch. f. Gynækol.*, Vol. 3, 1872.

⁴ Panum. *Virch. Arch.*, Vol. 60, 1874.

⁵ Metchnikoff. "L'Immunité dans les Maladies Infectieuses."

“L'exsudat inflammatoire doit être considéré comme une réaction contre toutes sortes de lésions et l'exsudation est un phénomène plus primitif et plus ancien que le rôle du système nerveux et des vaisseaux dans l'inflammation.”⁶

He compared the process of cell ingestion or phagocytosis of foreign particles, as here observed, to that taking place in the most simple intracellular digestion which occurs in unicellular forms, a hereditary cell function now specialized in certain mesodermal cells, and passed on in the evolution of higher forms to other specialized cells. And indeed in animals of the most complex structure the leukocytes which carry on this phagocytic process may be considered as, in a way, representing a primitive form of cell, since they are only nucleated elements of the body which wander from place to place, and are anatomically independent of nervous control. In 1883, at the Naturalists' Congress in Odessa, Metchnikoff⁷ first expressed his views and communicated the first of the splendid researches upon which our modern conception of phagocytosis is based.

His earlier studies were carried out with a small crustacean, the daphnia, in which he studied the reaction which followed the introduction of yeast cells. He observed the struggle which ensued between the ameboid leukocytes of the crustacean and the infecting agents and determined that complete enclosure of the yeast within the leukocytes assured protection to the daphnia, while a failure of this process, either from fortuitous causes or because of too large a quantity of the infecting agents, resulted in disease and rapid death.

This early work of Metchnikoff forms the beginning of a long train of investigations to which we owe most of the basic facts we possess concerning the rôle of the phagocytic cells in the protection of the body against infection. Just as the various serum phenomena, of which we have spoken, have a general biological significance apart from their importance in relation to bacterial invasion, so the process of phagocytosis must be looked upon as an attribute of the animal and vegetable cell which has important physiological bearing entirely apart from infection.

In fact, the ingestion of bacteria and other foreign particles by the leukocytes and other phagocytic cells of higher plants and animals is entirely analogous to the intracellular digestive processes which take place, as the ordinary manner of nutrition, among the unicellular forms. Among the rhyzopods, in general, food is taken in by means of the ingestion of other smaller forms of life, bacteria,

⁶ Inflammatory exudation should be considered as a reaction against all sorts of injuries, and exudation is a phenomenon more primitive and ancient than are the parts played by nervous system and blood vessels in the process of inflammation.

⁷ Metchnikoff. *Arb. a. d. zool. Inst.*, Wien, Vol. 5, 1883.

algæ, etc. (or particles of dead organic matter), into the cell body of the protozoön.

These materials are gradually engulfed by the body of the ameba, which flows about them with its pseudopods, and within the cytoplasm undergo gradual digestion. The process has been carefully studied by Mouton.⁸ In symbiotic cultures of amebæ with colon bacilli on agar plates, the bacteria are taken up in large numbers and about them are formed small vacuoles. That the digestion takes place in a slightly acid medium with the vacuoles can be proved by adding a drop of neutral red to the hang-drop preparation of amebæ and observing the brownish-red color taken by the materials in the vacuoles. Mouton was able to obtain a digestive ferment from the amebæ, by glycerin extraction, which exerted strong proteolytic action upon various albuminous substances, liquefied gelatin, and digested dead colon bacilli in vitro, acting best in slightly alkaline, but also in slightly acid, reactions. It is plain, therefore, that the most primitive form of digestion is an intracellular one carried on by ferments comparable in every way to the secreted digestive enzymes which accomplish the same purpose outside of the cells in higher animals. In essence, however, there is no fundamental difference physiologically between intra- and extracellular digestions, and the intracellular manner of assimilating solid nutritive particles may be retained in forms much higher in the scale of evolution than the rhyzopods. It has been studied by Metchnikoff and others in certain of the flat worms (*Dendrocelum lacteum*) in which typical phagocytosis is carried on by the cells of the intestinal mucosa. Many of these planaria obtain their nourishment by sucking the blood of higher animals. Placed under a microscope after feeding, it may be seen that the foreign blood cells are rapidly taken up by the intestinal epithelial cells, which engulf them by means of pseudopodia not unlike those of the ameba. After ingestion, here, too, the blood cells are surrounded by vacuoles within which their gradual disintegration or digestion is accomplished. Similar intracellular digestion seems to be general among the cœlenterates, and has been thoroughly studied by Metchnikoff in the actinia. Here the food particles are carried by the tentacles into the esophagus, and are taken up by the endodermal cells of the so-called "mesenteric filaments," where they are digested by a trypsin-like enzyme. In these animals digestion is entirely intracellular, though the ingesting cells are the parts of a specialized tissue. In other forms, still higher in the scale, although there is persistence of intracellular digestion, the extracellular process begins to be developed. Thus in certain mollusca the solid food is taken into the intestinal canal, where it first undergoes a preliminary digestion by secreted intestinal juices. After it has

⁸ Mouton. *C. R. de l'Acad. des Sciences*, Vol. 133, 1901.

been reduced to small amorphous particles in this way, these are seized by the ameboid cells, and intracellular digestion completes the process which has been begun extracellularly.

As we study the process among higher animals, it appears that, among vertebrates, the intracellular methods of digestion have been, at least for normal metabolism, entirely displaced by the extracellular as it occurs in the intestine, where solid particles are rendered completely amorphous, dissolved, and reduced to a diffusible condition by the digestive juices before they are offered to the cells for utilization. However, the capacity for intracellular digestion is not entirely lost, and is retained of necessity in certain body cells. For were there not such an emergency arrangement the body would lack an available mechanism with which to meet such accidents as extravasations of blood, or the entrance of bacteria and other foreign solid particles into the tissues. It seems reasonable to classify both the phagocytic action of body cells and the formation of antibodies in the blood plasma, primarily as emergency devices for the digestion of foreign materials both formed and unformed which, under abnormal conditions, penetrate into the physiological interior of the body (blood stream or tissue spaces), and must be disposed of.

In the lowest animals the single cell is called upon to perform all necessary functions. In the course of evolution, however, as the body becomes more and more a community of many cells, a division of labor takes place which is expressed morphologically in the differentiation of tissues and organs, and physiologically in the adaptation of individual tissue cells to the performance of specialized functions. Nevertheless, it is necessary, both for certain normal processes, as well as for provision against such complex emergencies as those mentioned, that certain cells of the complex community should retain the primitive abilities of the more independent cells of the lower forms. Thus, among many animals, the phagocytic action of cells performs definite services in the course of normal development. This is seen most markedly in some insects (diptera) in which the destruction of larval organs, useless to the adult animal, may be entirely accomplished by the action of phagocytic cells, and a similar process may accompany the transformation of the tadpole to the adult in many amphibia.⁹ In higher animals the removal of extravasations of blood is accompanied by a train of occurrences which is readily subjected to study.¹⁰ In such cases the leukocytes rapidly enter the area of extravasation and an engulfment of the blood cells occurs, followed by a process of digestion entirely analogous to the digestion of similar blood elements by the various forms of intestinal hemamebæ. In the latter case it is a process of normal digestion, in the

⁹ See Henneguy. "Les Insectes," Paris, 1904, p. 677.

¹⁰ Langhans. *Virchow's Archiv*, Vol. 49, 1870.

former an emergency procedure carried out by virtue of the retained ancestral characteristics of the special phagocytic cells.

The leukocytes, whose chief functions seem to be associated with such processes of intracellular digestion, may, therefore, be looked upon as cells retaining primitive characteristics for definite physiological purposes. We shall see, however, that, to meet exceptional conditions, the process of phagocytosis may be carried out also by many other cells which are associated ordinarily with functions entirely apart from this phenomenon.

During normal life in higher animals, too, constant destruction of red blood cells by phagocytosis takes place in the spleen and liver, and is described by Dickson¹¹ as occurring in the bone marrow as well; and similar phagocytosis of red cells is seen in the hemolymph nodes. It is claimed by Metchnikoff, furthermore, that many of the degenerative and retrogressive processes which take place in the human body are carried on by the mechanism of phagocytosis. The rapid return of the puerperal uterus to the normal state is explained in this way, and work by Helme¹² seems to show that there is an actual phagocytosis of the hyperplastic uterine musculature during this period. The atrophic changes of senility, too, are attributed by Metchnikoff^{13 14} to the same processes. The involution of the ovaries is accompanied by active phagocytosis of portions of this organ, and Metchnikoff claims further to have shown that the degeneration of the nervous system during old age is accomplished by the phagocytosis of nerve cells by phagocytic elements derived either from the leukocytes or the neuroglia, or from both.¹⁵ The whitening of the hair, both in human beings and in old animals (dogs), is similarly due, he claims, to phagocytosis of the pigment by cells which wander in from the root sheaths. It is, up to the present time, impossible to determine the stimulus to which this phagocytosis is due.

Since the subject is a very important one, many studies have been made to determine which cells of the body of higher animals can take in and digest foreign particles and to classify them according to this power. Metchnikoff has distinguished between the "motile" and "fixed" phagocytes, the former the leukocytes of the circulating blood, the latter certain connective tissue cells, endothelial cells, splenic pulp cells, and certain cellular elements of the lymph nodes,

¹¹ Dickson. "The Bone Marrow," Longmans, Green, London, 1908.

¹² Helme. *Transact. Roy. Soc. of Edinburgh*, Vol. 35, 1889. Cited from Metchnikoff.

¹³ Matschinsky. *Ann. de l'Inst. Past.*, Vol. 14, 1900.

¹⁴ Metchnikoff. *Ann. de l'Inst. Past.*, Vol. 15, 1901.

¹⁵ That the leukocytes are concerned in the destruction and resorption of dead tissues has been shown by Leber especially (Leber, "Die Entstehung der Entzündung," Leipzig, Engelmann, 1891). An accumulation of leukocytes about a bacterial focus or from any other stimulus is followed by tissue lysis due to leukocytic enzymes.

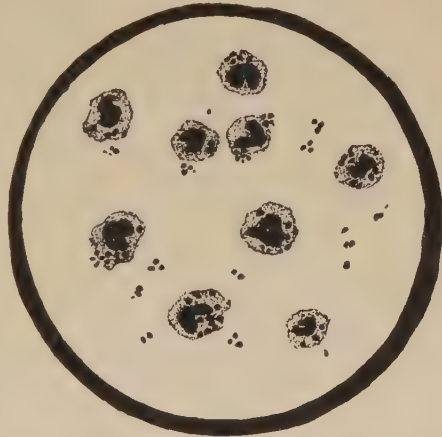
the neuroglia tissue, and, in fact, all phagocytic cells which are ordinarily confined to some definite localization in the body. Among phagocytic cells Metchnikoff further distinguishes between "microphages," by which he designates the polymorphonuclear leukocytes of the circulating blood and "macrophages." The macrophages include the fixed cells mentioned above, together with the large mononuclear elements of the blood, in short; all phagocytic cells except the microphages.

Although no absolute functional differentiation is possible between the two, it is true, in a general way, that the microphages are concerned primarily with the phagocytosis of bacteria and especially of those which invade acutely, while the macrophages are con-

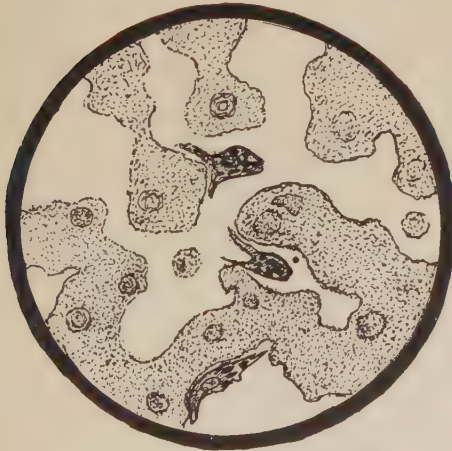
cerned especially with the resorption of cellular detritus, foreign bodies, and such bacteria as are more chronic in their activities, or are peculiarly insoluble.

On the other hand, microphages may take up foreign particles and bacteria of all kinds under suitable conditions, and no sharp line can be drawn between the two varieties in this respect. Metchnikoff further believes that the two classes of phagocytic cells differ in the nature of the protective substances they secrete and furnish in the blood plasma. This, however, is a problem concerning which there is much difference of opinion and which calls for separate discussion in another place.

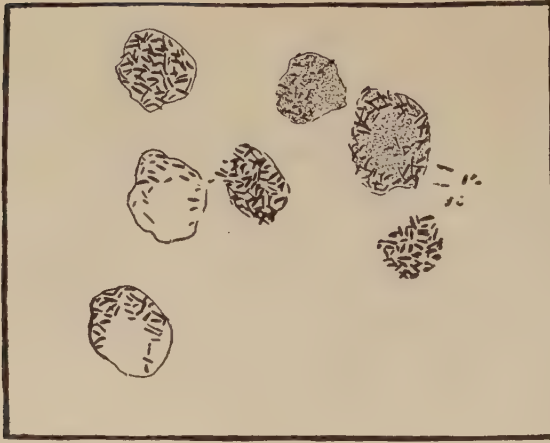
The property of phago-



POLYNUCLEAR LEUKOCYTES TAKING UP STAPHYLOCOCCI.



KUPFER CELLS CONTAINING MALARIAL PIGMENT. DIAGRAMMATICALLY DRAWN FROM A SECTION OF MALARIAL LIVER KINDLY FURNISHED BY DR. R. LAMBERT.

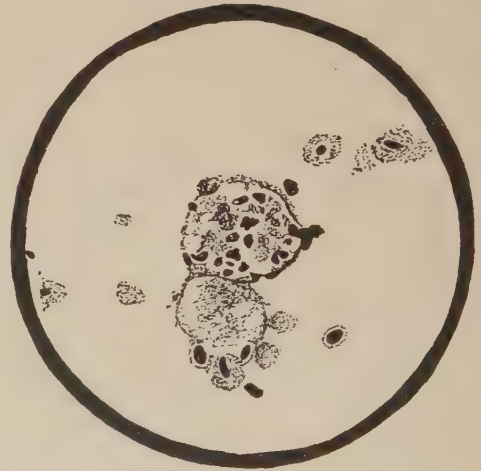


RAT LEPROSY BACILLI GROUPED IN THE REMAINS OF DEAD SPLEEN CELLS GROWING IN PLASMA.

Drawn after illustration in Zinsser and Carey, *Journal of the A. M. A.*, Vol. 58, 1912.

cytosis is therefore an attribute of a considerable number of different varieties of cells. In the circulating blood the polynuclear leukocytes are the most actively motile and phagocytic elements. The eosinophile cells may also take up foreign particles and bacteria, as may also the large lymphocytes. The small lymphocytes and mast cells are either entirely inac-

tive in this respect, or, at least, possess phagocytic powers under exceptional circumstances only. This does not mean, however, that these last-named cells may not accumulate at the point of invasion nor that they may not play an important part in the defence of the body. It is well-known, of course, that, in tuberculosis and a number of other conditions, the lymphocytes may form the majority of the cellular elements which accumulate at the site of the lesion. Among the fixed cells of the body it is probable that phagocytosis may be carried on by cells of many different origins, though the identification of cells in tissues is often a purely morphological problem, and therefore fraught with many possibilities of error. Probably the most active fixed tissue cells are the endothelial cells of the blood vessels and those which line the serous cavities, the



PHAGOCYTOSIS OF SENSITIZED PIGEON CORPUSCLES BY ALVEOLAR CELLS OF LUNG.

Drawing made after photomicrograph published by Briscoe, *Journal of Path. and Bact.*, Vol. 12, 1908.

sinuses of the lymphnodes, and of the spleen. However, there are many other cells in addition to these which may be phagocytic. The writer, with Carey,¹⁶ has observed the active phagocytosis of leprosy bacilli by cells, probably of connective tissue origin, growing from plants of rat spleen in plasma. Phagocytosis by the cells lining the alveoli of the lungs has been observed by Briscoe.¹⁷ This author made the interesting observation that in cases of mild infection such cells can free the lungs of micro-organisms entirely without aid from the leukocytes of the circulating blood. It is these cells, too, which, in the ordinary conditions of life, take up the inhaled particles of dust and are, therefore, often spoken of as dust cells. The origin of the dust cells has often been the subject of controversy. In the embryo the alveoli of the lung, like the bronchi, are lined with columnar cells which are transformed into flattened epithelium as the alveoli expand at the first inspirations after birth. These flattened cells, which constitute the alveolar or dust cells, are probably of epithelial origin, and as such are probably the only epithelial cells which act as phagocytes under ordinary conditions. Although no positive general statement is justified, we can yet say with reasonable accuracy that among the phagocytic fixed tissue cells the most important are the connective tissue and endothelial cells.

The type of phagocytosis and the variety of cell which participates in it seem to depend to a great extent upon the nature of the substance which incites the process, or rather at which the process is aimed. Thus the large cells which, in tissues, take up the leprosy bacillus, those which are characteristic of tuberculous foci, or those caused by blastomycetes, or by foreign bodies, all have special appearances which are sufficiently characteristic to have diagnostic value.

However, it is difficult to determine with certainty the origin of the cells which participate. The chemical nature of the substances taken up, moreover, often complicates the phagocytic process in such a way that different cellular elements are enlisted in succession in order that the ingested substances may be disposed of. Thus tubercle



GIANT CELL IN TUBERCULOSIS.

¹⁶ Zinsser and Carey. *Jour. A. M. A.*, March, 1912, Vol. 58.

¹⁷ Briscoe. *Jour. of Path. and Bacter.*, Vol. 12, 1907.

or leprosy bacilli which are injected into an animal may be at first taken up by polynuclear leukocytes or microphages, by which they may even be carried into the lymph channels and distributed, perhaps to the detriment of the host. But these cells, probably because they lack a lipolytic ferment by means of which the waxes of the acid-fast organisms can be digested, cannot destroy the bacteria, which are then attacked by other cellular elements at the site of their final deposit.

In many such cases the further resolution of the foreign substance is accomplished by an important type of phagocytosis which is characterized by the formation of the so-called giant cells. These cells are of varying appearance in different conditions and locations. Thus the giant cells which form about foreign bodies, such as the



FOREIGN BODY OF GIANT CELL. SECTION OF CORNEA AFTER EXPERIMENTAL INJECTION OF PARAFFIN.

After preparation kindly furnished by Dr. W. C. Clarke.

small cotton fibers occasionally left in wounds, or injected particles of paraffin or iron splinters, etc., are quite characteristic and distinct from the giant cells of tuberculous foci, or of rhinoscleroma, glanders, or leprosy. They are all large cells, containing often numerous nuclei which form either by the fusion of several cells, as claimed by Borrell,¹⁸ Hektoen,¹⁹ and others, or by the cleavage of the nuclei alone, without coincident divisions of the cytoplasm.

Although it is, of course, impossible to decide definitely upon purely morphological grounds, the

researches of Hektoen especially would lead one strongly to favor the former view. It is equally difficult to decide the origin of giant cells, and endothelial, connective tissue, and even leukocytic origin has been claimed for them. Yet in no case has it thus far been possible to actually observe their formation by a method which could positively decide this point.

In order to gain a clear conception of the participation of phagocytes in the response of the body to injury or invasion, it will be useful to follow out the process of inflammation as it occurs in the

¹⁸ Borrell. *Ann. de l'Inst. Past.*, 7, 1893.

¹⁹ Hektoen. *Jour. Exp. Med.*, 3, 1898, p. 21.

higher animals. Inflammation may be incited by a large number of agencies—chemical irritants, mechanical injury, or even by the introduction of inactive and isotonic substances such as broth or salt solution.^{20 21} Yet in these cases the response, though essentially similar in principle to that following invasion by bacteria, lacks certain features especially interesting in the present connection, and it will be most profitable for our purpose to consider in detail the result of infection with pathogenic micro-organisms.

If an emulsion of pyogenic staphylococci is injected into an animal subcutaneously the site of injection will soon become reddened and swollen and microscopic examination will show, within a few hours, a swelling and engorgement of the blood vessels.

The injected cocci will be found to lie partly scattered in the tissue spaces, in part within polynuclear leukocytes and connective tissue cells which have begun to ingest them. The tissue spaces will be swollen and stretched by the exudation of blood serum from the vessels. This condition will begin in from 4 to 6 hours after injection and increase during the next 24 hours in extent and severity, according to the quantity and virulence of the cocci injected. The conditions which precede the wandering of the polymorphonuclear leukocytes out of the vessels have been carefully studied in such thin tissues as the mesentery of a frog after injury by trauma or acid. Within the vessels of the affected area there is at first an acceleration of the blood stream, then a dilatation of the capillaries and a slowing of the current. Leukocytes may now be observed moving more slowly than the main stream, and keeping close to the periphery along the walls of the vessels. Here and there they seem interrupted in their movements and adhere to the vascular wall. A little later these cells appear to pass through the wall of the



DIAGRAMMATIC REPRESENTATION OF LEUKOCYTES WANDERING THROUGH CAPILLARY WALLS.

Adapted from Ribbert, "Lehrbuch der Allgemeinen Pathologie," p. 337.

injury by trauma or acid. Within the vessels of the affected area there is at first an acceleration of the blood stream, then a dilatation of the capillaries and a slowing of the current. Leukocytes may now be observed moving more slowly than the main stream, and keeping close to the periphery along the walls of the vessels. Here and there they seem interrupted in their movements and adhere to the vascular wall. A little later these cells appear to pass through the wall of the

²⁰ See Adami. "Inflammation," Macmillan, London, 1909.

²¹ See Adami, *loc. cit.*

vessel by sending out pseudopodia which slowly penetrate it. Adami states that if, at this stage, the tissues be excised, fixed in osmic acid, and stained, leukocytes may be seen crowding the inner surface of the vessel in all stages of transition from its anterior to the lymph spaces on the outside.

In the staphylococcus infection, after from 12 to 48 hours, there will be seen the results of an active and destructive struggle between the invading bacteria and the defending cells. In the center of the area of invasion tissue has been destroyed and disintegrated. Amid the necrotic detritus, closely packed, lie leukocytes and cocci and active phagocytosis has taken place. In some cases the intracellular bacteria appear swollen and disintegrating, in others the leukocyte itself, overcome by the larger number of bacteria it has taken in, becomes vacuolated, indefinite in outline, and apparently is being itself destroyed. The presence of blood serum, which is aiding in the destruction of bacteria both by its bactericidal powers and its reinforcement of the phagocytic process, renders this mass fluid or semi-fluid, and the whole mixture constitutes what is known as pus. Around the periphery cocci and leukocytes become more scattered and sparse, and bacteria, together with leukocytes, loaded with cocci, may be seen lying within large mononuclear cells (macrophages). Whether the process goes on to further extension or is eventually walled off into a distinct abscess by the formation of granulation tissue and new connective tissue depends upon the balance of forces between attacking agent and defensive factors.

If we inject a similar emulsion of cocci into the pleural or peritoneal cavity of an animal a process similar in principle may be observed.

Normally the peritoneum contains a small amount of this serous fluid and a moderate number of white blood cells, chiefly lymphocytes. When any substance, broth or salt solution, an aleuronat or a bacterial emulsion, is injected into the peritoneal cavity, there follows a brief period during which there is a diminution of the free cellular elements in the peritoneal fluid. At this time there is a clumping of cells in the folds of the omentum and mesentery, a transient stage of flight away from the point of injury. This, however, is soon over. Within one to two hours an active immigration of leukocytes into the serous cavity occurs and if, during the next 12 to 24 hours small quantities of fluid are, from time to time, withdrawn with a capillary pipette, a rapid and constant increase of leukocytic elements, chiefly of the microphage or polynuclear type, is observed. If the injected substance has been a sterile, harmless fluid, a gradual return to normal within 48 hours then ensues. If, however, we have injected bacteria, a struggle similar to the one described above takes place within the peritoneum, and active phagocytosis of the micro-organisms takes place.

Let us suppose that the injected bacteria have been small in quantity and moderate in virulence. In such a case a rapid phagocytosis gradually rids the fluid of micro-organisms and within 24 hours after injection few, if any, free bacteria are visible.

A little exudate taken at this time shows large numbers of microphages varyingly crowded with well-preserved and disintegrating bacteria. Some of the phagocytes, having literally taken up more than they can digest, are vacuolated and disintegrating, but, in general, the victory lies with the cells. A little later large mononuclear elements appear, and here and there will be seen to take up dead leukocytes together with ingested cocci. In this way gradually a cleaning out of the peritoneum takes place, the animal recovers, and the peritoneum returns to normal.

Let us suppose, on the other hand, that the bacteria injected are in larger doses and of greater virulence. In such a case, after a period of active phagocytosis, there may be a gradual increase of bacteria over leukocytes. The phagocytic cells are found to be undergoing degeneration in larger numbers, the free bacteria increase, and the impending death of the animal can often be foretold by the appearance of the exudate. Finally, the peritoneal fluid may consist chiefly of free and rapidly multiplying bacteria with a practical absence of phagocytic cells.

In all of the processes so far as described the burden of the defence has fallen upon the microphages or polynuclear leukocytes, while the macrophages—endothelial and connective tissue cells—have taken a purely secondary part in the reaction, forming, to some extent, a second line of defence, or, more probably, taking part only in the final removal of degenerated and disintegrating combatants and tissue detritus. In order to obtain a complete conception of phagocytosis in its entire significance it will be necessary to consider a further example, namely, the process which takes place within tissues in the course of the efforts of macrophages to remove bacteria and other substances which, either because of their insolubility or for other unknown reasons, are refractory to the attacks of the microphages. Since we are interested in this subject chiefly from the point of view of the defence against bacteria, we may illustrate this process best by the description of the reaction which takes place when tubercle bacilli become localized anywhere within the animal body.

When tubercle bacilli are injected into the peritoneum they are actively taken up by the polynuclear leukocytes just as are other bacteria and many entirely inactive solid particles. A similar ingestion by microphages may take place in the folds of the intestinal mucosa if tubercle bacilli are fed to guinea pigs. However, this preliminary phagocytosis is probably of but secondary significance in the combat of the body against tuberculosis, since it has still to be shown that polynuclear leukocytes are capable of digesting and

destroying acid-fast bacilli. Indeed, much evidence tends to show that the ingestion of tubercle bacilli by microphages may be a detriment to the host, since the bacilli by this means are carried through the lymphatics and variously distributed throughout the body. Polynuclear leukocyte extracts, though containing, as we shall see, proteolytic enzymes, do not, according to Tschernorutzky, contain any lipase, and it may well be that for this reason they are unable to attack the waxy substances which form an integral part of these organisms. This is in keeping with the observations made by Terry in our laboratory, that rat leprosy bacilli may be kept within leukocytes for weeks without losing their acid-fast properties, whereas the same bacilli, as the writer and Cary found, were rapidly disintegrated in spleen cells growing in plasma. Moreover, it is well known that the estimation of tuberculo-opsonin contents of the sera of tuberculous patients has been peculiarly unsatisfactory in throwing light on the progress of the disease. It would seem, therefore, that in this disease, as well as in others caused by acid-fast organisms, the microphages play only an unimportant part in the defence of the body.

On the other hand, when tubercle bacilli are deposited either in a lymphnode (through the vehicle of leukocytes) or in a capillary anywhere by the blood stream, a train of cellular changes is initiated in which the predominant part is played by the macrophages. The tubercle bacilli so deposited are rapidly surrounded by large mononuclear cells, probably endothelial in origin. Some of the microorganisms may even be phagocyted and taken into these cells. These cells, spoken of as "epithelioid cells," surround the clump of bacteria in more or less concentric rings, and around these there is an accumulation of leukocytes, largely of the lymphocyte variety, with an admixture of a very few microphages. Then by the fusion of endothelial cells, or possibly by division of the nuclei of some of these cells within the individual cell bodies, giant cells are formed which take up the bacilli. The further progress of the tubercle now greatly depends upon the balance of power. Often such a tubercle may heal, possibly because of complete intracellular digestion of the bacilli. On the other hand growth and multiplication may lead to a slow and dry necrosis of the center of such a mass of cells, leading to the condition spoken of as caseation. Epithelioid cells lose their outlines and staining properties, and go to pieces. The center of the lesion is a grumous mass, the periphery shows a few giant cells and connective tissue proliferation.

It is always surprising to those who study these lesions for the first time how rarely they succeed in finding tubercle bacilli in microscopic sections prepared from such tubercles by the ordinary Ziehl-Neelsen method of staining. Repeated and careful examination of such material may fail to reveal any acid-fast organisms,

though inoculation into guinea pigs is nevertheless successful, producing typical tuberculosis. Much²² has studied this peculiar state of affairs particularly and has shown that, although such lesions may show no tubercle bacilli by the Ziehl-Neelsen carbol-fuchsin method, staining by a modified Gram technique will reveal numerous Gram-positive rods and granules which have lost their acid-fast properties. This, too, if true, and the evidence is very much in its favor, would point to an ability of the macrophages to digest the waxy substance of the tubercle and other acid-fast bacilli, a property not possessed by the microphages. It may, of course, mean on the other hand that the tubercle bacilli in the lesion have not developed the waxy condition.

CHEMOTAXIS AND LEUKOCYTOSIS

The part played by the phagocytic cell in the defence of the body against the entrance of bacteria and other foreign substances consists, then, of two functionally different phases. The first is an active motion of the cells toward the point attacked, and their accumulation about the noxious agent, the second consists in the act of ingestion itself.

The motion of the leukocytes toward the invading substances indicates a sensibility on the part of the cell to changes in its environment incited by the foreign agent, and since the stimuli most likely to reach the leukocytes and bring about this alteration in the direction of their movements are chemical in nature, the phenomenon is spoken of as "chemotaxis." This term was borrowed from Pfeffer,²³ who studied similar phenomena in connection with many freely motile plant cells, spermatozoa, and bacteria. Since the change of direction brought about in a moving cell by such influences may be such as either to attract or to repel, the term "positive chemotaxis" is used to designate the former and that of "negative chemotaxis" the latter.

The property of chemotaxis is of vital interest in the present connection, since, whatever may be our opinion regarding the relative values of phagocytosis and serum protection in immunity, the great importance of the phagocytic process cannot be questioned, and any agency which repels the approach of the phagocytes must be a detriment, while any factor which attracts them is, of necessity, a powerful means of defence. In the investigations upon the nature of infectious diseases attention has been concentrated upon the phenomenon of phagocytosis, and the relations governing the act of ingestion have been very thoroughly studied. The details of the chemotactic phenomenon, however, though of equal importance, are much more

²² Much. "Beiträge zur Klinik der Tuberk.," Vol. 8, 1907, Hft. 1 and 4.

²³ Pfeffer. "Untersuch. a. d. Botan. Inst. Tübingen," Vols. 1 and 2, 1884 and 1888.

obscure. A large part of our sparse knowledge in this connection, moreover, has been gained by studies not related to infection.

The stimuli which determine the motion of cells are, of course, not necessarily chemical, and extensive studies have been made upon the effect of light waves in this connection. Although these investigations are of great biological importance, they have little direct bearing upon the problems of tropism as related to bacteria and leucocytes and cannot therefore be considered here.

Some of the earlier researches upon chemotaxis were those made by Stahl²⁴ upon the slime-molds or myxomycetes. These organisms possess the power of ameboid motion, and were observed by Stahl to move toward or away from any given region, according to the nature of the substances with which they came in contact. Pfeffer subjected this phenomenon to closer analysis. Working with the spermatozoa of ferns, swarm spores, bacteria and infusoria, he elaborated an ingenious technique by means of which he was enabled to determine directly the negative or positive chemotactic properties of various substances in solution upon these motile forms. His technique was exceedingly simple. Capillary glass tubes, about 8 to 10 mm. long and 0.1 mm. in diameter, were sealed at one end in the flame, and then dropped into a watch-glass. The solution which was to be tested was poured over the tubes and the watch-glass then placed under the bell of an air-pump. When the air was evacuated and pressure reduced the tubes became partly filled up with the liquid. They were then removed, washed in water, and placed under a cover slip under which a preparation of the motile cells was swimming. Positive chemotaxis was indicated by entrance of the cells into the tubes, negative, by their refusal to enter. Failure of the solution to exert any chemotactic influence resulted in their moving into and out of the tubes indiscriminately.²⁵

By this technique a large number of interesting observations were made which threw much light upon the causes underlying the movements of plant cells. For instance, in investigating the spermatozoa of the ferns it was found that they were attracted strongly by malic acid and its salts, while no other substance investigated approached these compounds in the intensity of positively chemotactic stimulation. From this Pfeffer concludes that the bursting of the fern archegonia is accompanied by the liberation of malic acid, this attracting the male to the female cell.

Similar experiments have been carried out since then by numerous naturalists, among them Buller,²⁶ Lidforss,²⁷ and Jennings,²⁸

²⁴ Stahl. *Botanische Zeitung*, 1884.

²⁵ Buller. *Annals of Botany*, Vol. 16, No. 56, 1900.

²⁶ Buller. *Loc. cit.*

²⁷ Lidforss. "Jahrbücher f. wissensch. Botanik," 41, 1904.

²⁸ Jennings. "Behavior of Lower Organisms," Columbia Univ. Press, Macmillan, 1906.

and it has been found that in addition to malic acid compounds many other substances, organic and inorganic, occurring in plant cells and cell-sap exert positive chemotactic power. Lidforss has shown, for instance, that calcium chlorid in 0.1 per cent. solution may strongly attract plant spermatozoids (equisetum—horsetail). When the solution is concentrated to 1 per cent., attraction is still exerted, but the spermatozoids immediately lose their motility upon entrance into the fluid.

The same worker has shown that a substance which is positively chemotactic for one variety of plant cell may be negatively chemotactic for another, showing a certain selective variation which should be of great biological importance. Thus capillaries with a 1 per cent. solution of potassium malate actively attracted the spermatozoids of marchantia (a liverwort), while not a single spermatozoid of equisetum would enter these tubes. Löw²⁹ has applied these methods of study to the investigation of the chemotaxis of mammalian spermatozoa and found that these cells were actively attracted by weakly alkaline solutions.

Studies upon the factors determining the movement of bacteria and amebæ toward some substances and away from others have been numerous, and are valuable for the understanding of leukocytic chemotaxis, because they have led to the formulation of a number of important general theories. The fact that the motions of bacteria in suspensions are, to a certain extent, determined by the negative electrical charge which they all carry in neutral media, has been touched upon in the section on agglutination. Attempts on the part of Young and the writer to determine whether the attraction of leukocytes toward bacteria might be due to the carrying of an electropositive charge by the white cells have met with no result, owing so far to the failure to elaborate a reliable technique. However, this thought is not an impossible one and should be borne in mind.

That certain bacteria will wander actively toward a source of oxygen was shown by Engelmann's³⁰ classical experiment in which a diatom, half in the shade and half in the light, was surrounded by an emulsion of bacteria, and these were seen to collect about the lighted half only, where oxygen was being liberated by virtue of the chlorophyll. The extreme delicacy of chemotactic reactions is illustrated in these experiments in that Engelmann calculated that the bacteria reacted to one one-hundred billionth of a milligram of oxygen. The selective reaction of bacteria to various chemical substances, furthermore, has been shown by allowing different solutions to diffuse into bacterial emulsions from capillary tubes, and by observing attraction or repulsion from the point of contact.

The chemotaxis of leukocytes has opposed more difficulties to

²⁹ Löw. *Sitzungs Berichte kais. Akad. d. Wiss.*, Wien, Vol. 3, Abf. 3.

³⁰ Engelmann. *Arch. f. d. ges. Physiol.*, Vol. 57, p. 375.

direct study, since the conditions within the living body are subject to a large number of modifying factors, and experiments upon the isolated cells, in vitro, even under conditions of the most careful technique, are fraught with much unavoidable injury to the cells. However, enough has been learned to indicate that these cells are subject to the phenomena of chemotaxis or tropism just as are independent unicellular forms, and that they may be attracted or repelled by a variety of organic and inorganic substances. Leber³¹ was one of the first to study this in his work upon inflammation. He found that leukocytes were actively attracted by powdered copper and mercury compounds, but not by powdered gold or iron. He also observed that dead bacteria exerted a similar positive chemotactic influence, and Buchner³² later succeeded in extracting substances from various bacteria which possessed similar properties. It appears, from these and other investigations, that the power of stimulating positive chemotaxis is a general property of bacterial proteins, equally evident in bacterial extracts, dead bacteria, or the living organisms. It is likely, therefore, that the attraction of leukocytes toward the point of bacterial invasion is, in part at least, due to the properties of the bacterial proteins themselves. That this, however, is not the whole story is evident from the work of Massart and Bordet,³³ who showed that the products of cell destruction and disintegration possess similar positively chemotactic properties. This is true not only of the products of disintegrated tissue cells, but of those of the destroyed leukocytes themselves. Thus it appears that when any injury of tissue takes place, a stimulus which attracts leukocytes results, even when the injury is not accompanied by bacterial invasion. This would explain the participation of leukocytes in reactions to injury, and in inflammations not of bacterial origin, and their local accumulation following the injection of insoluble inorganic substances.

When bacteria are actually present, however, the added stimulus due to the diffusion of bacterial proteins probably increases the process to a degree often sufficient to meet the added requirements for protection. Following this, both the destruction of tissues, of bacteria, and of leukocytes may together exert a cumulative chemotactic power which continues the process proportionately with the extent of the lesion.

It is of the utmost importance, therefore, to ascertain whether or not any substances derived from bacteria may, under any circumstances, exert a repellent or negatively chemotactic power. If we infect an animal intraperitoneally with virulent bacteria, in doses

³¹ Leber. *Fortschr. der Med.*, 1888; also "Die Entstehung der Entzündung," Engelmann, Leipzig, 1891.

³² Buchner. *Berl. klin. Woch.*, Vol. 27, No. 30, 1890.

³³ Massart and Bordet. *Ann. de l'Inst. Past.*, Vol. 5, 1891.

sufficient to lead to death, and examine the peritoneal exudate just before the lethal outcome, we may observe that leukocytes are gradually disappearing, and that finally but a few will be present and the fluid will be swimming with free micro-organisms. In the same way it is well known that the diminution of leukocytes in the circulating blood—or even the failure of these cells to increase in the circulation in the course of such diseases as pneumonia, or general infections with staphylococci or streptococci—is seriously prognostic of fatal outcome. The conditions here observed point strongly to the existence of substances of negative chemotactic influence which protect the bacteria, not from phagocytosis itself, but from that necessary forerunner of phagocytosis, the approach of the leukocyte. It is necessary to draw this distinction since these phenomena are not merely, as often believed, “antiopsonic,” but in truth largely “anti-chemotactic.” It is true that Kanthack,³⁴ and more especially Werigo,³⁵ have denied the existence of negatively chemotactic bacterial products, the latter basing his assertion upon the observation that active phagocytosis occurs in the lungs, liver, and spleen of animals dying of infection with virulent germs. However, the arguments of these authors are not conclusive and the mass of experimental and clinical evidence which points to a direct failure of leukocyte accumulation in the presence of virulent bacteria in the animal body would alone suffice to render such conclusions unlikely. Moreover, strong evidence in favor of the existence of negatively chemotactic influences is brought by the extensive experiments of Bail upon the so-called aggressins, discussed in another place, and such observations as those of Vaillard and Vincent³⁶ and Vaillard and Rouget,³⁷ which showed that the injection of a little tetanus toxin together with tetanus spores would prevent the ingestion of the spores by leukocytes, and thereby furnish an opportunity for germination and consequent fatal toxemia.

Similar observations have been made by Besson³⁸ in the case of the bacillus of malignant edema by the use of the original technique of Pfeiffer. Capillary tubes containing the toxin remained free of leukocytes after subcutaneous introduction into guinea pigs, while similar tubes containing the culture medium alone, or the bacilli and their spores, attracted leukocytes in considerable numbers.

It is possible, of course, to interpret such phenomena as due to a failure of positive chemotaxis rather than to an active negative chemotaxis.

Although the phenomena of chemotaxis are most easily studied

³⁴ Kanthack. Quoted from Adami, *loc. cit.*

³⁵ Werigo. *Ann. de l'Inst. Past.*, Vol. 8, 1894.

³⁶ Vaillard and Vincent. *Ann. de l'Inst. Past.*, Vol. 5, 1891.

³⁷ Vaillard and Rouget. *Ann. de l'Inst. Past.*, Vol. 6, 1892.

³⁸ Besson. *Ann. de l'Inst. Past.*, Vol. 9, 1895.

in extravascular inflammatory changes, there is none the less a regular and apparently purposeful attraction or repulsion of leukocytes evident in the circulating blood during infectious diseases. That infection of the body with many micro-organisms results in the increase of leukocytes, and that in others there is either no increase or even a decrease, is too well known and too generally applied in diagnosis and prognosis to warrant our giving up much space to a review of the facts. Nevertheless, the causes which lead to a leukocytosis in the one case, a leukopenia in the other, are still very obscure and deserve discussion.

In the first place it is by no means certain whether a leukocytosis signifies an active discharge of new leukocytes from the bone marrow or whether it means simply an altered distribution in that the phagocytes accumulated in the lymphatic and other organs are attracted by chemotaxis into the peripheral circulation. Studies of the bone marrow during infection as well as the occasional appearance of myelocytes and other cells ordinarily found only in the bone marrow during health would point toward a participation of active bone-marrow hyperplasia in the increase of peripheral leukocytes. There is no good reason to doubt, moreover, that a chemotactic stimulus exercised in the circulation should withdraw leukocytes from any place of accumulation to the circulation. Probably both processes take part. When bacteria are injected into the circulation of an animal there is, at first, a moderate diminution of the leukocytes just as there is after injection of bacteria or other substances into the peritoneum. This is soon followed in most cases by a rapid and progressive increase, in which, whenever the leukocytosis is one of considerable degree, the polynuclear leukocytes preponderate. The extensive clinical study of the white cells in infectious disease of the human being give us more material for reasoning in this respect than we have available from animal experiment. Infection with invasive bacteria such as the pneumococcus (and Neufeld and others have shown that most lobar pneumonias are accompanied by pneumococcus bacteriemia), streptococci, staphylococci, and others is always accompanied by an increase of the leukocytes, while, in typhoid fever, influenzal infection, tuberculosis, and a number of other infections, the leukocytes do not increase and may even decrease. How are we going to account for this? That all these bacteria contain a substance which is positive in its chemotactic effects is easily demonstrated by injecting them into the peritoneum and observing an accumulation of leukocytes and a consequent phagocytosis, even in the cases of those organisms which do not call forth a leukocytosis in the blood of the diseased human being. Thus it has been our experience as well as that of others invariably to observe the rapid and complete polynuclear phagocytosis of both leprosy bacilli and tubercle bacilli after injection of these micro-

organisms into the peritoneal cavities of guinea pigs. Yet a chronic tuberculous peritonitis or pleurisy is characterized usually by an exudate which contains but few polynuclears and relatively many lymphocytes. A final explanation of these conditions is not possible at present. No adequate explanation for the selective accumulation of lymphocytes and the absence of polynuclear cells about tuberculous foci has yet been advanced. The absence of polynuclear leukocytosis may possibly be due to the great insolubility of these bacilli, in consequence of which little or no leukocytosis-stimulating substances are liberated.

Pearce³⁹ has suggested a similar reason for the absence of polynuclear accumulations about chronic localized lesions of any kind in which tissue encapsulation may prevent the contact of the inciting agents with the body fluids and there is a consequently slow or slight production of such chemotactic stimulating materials.

In typhoid fever, where the slight primary leukocytosis is rapidly succeeded by a leukopenia with a relative lymphocytosis, the conditions are somewhat different. Here, as in some other infections, as Friedberger and others have shown, we are dealing with a generalized infection by an organism which is easily subject to the action of alexin with consequent production of anaphylatoxin. (See chapter on Anaphylaxis.) This poison, it seems, exerts a negative chemotaxis, and probably during the height of the disease, therefore, leads to the low leukocyte count observed. That this is at least likely seems to follow from the studies which have been made upon the nature of the typhoid poisons, and also from the observation of Gay and Claypole, that typhoid-immune rabbits react to the infection of typhoid bacilli with a rapid and powerful increase in the polynuclear leukocytes, whereas similar injections into the normal animal lead to leukopenia.

If the supposition regarding tuberculosis, made above, is correct, it would follow that a sudden and considerable increase in the polynuclear leukocytes in a case of tuberculosis would indicate a discharge of organisms into the circulation and a tendency toward generalization of the infection in this manner. (See Weigert's view of the manner in which tuberculosis may spread by the destruction of the wall of a vein by a localized lesion.) However, although speculation in the absence of experimental proof is justified, it must not be forgotten that the problems of selective chemotaxis are too obscure to permit of our laying much weight on any of these views.

Gabritchewsky,⁴⁰ who investigated this subject extensively, has classified various substances according to their positively, negatively, or neutral chemotactic activities. It is not necessary to recapitulate these, but it is interesting to note that he found that some substances

³⁹ Pearce. *Jour. A. M. A.*, Vol. 61, 1913.

⁴⁰ Gabritchewsky. *Ann. Past.*, Vol. 4, 1890.

which were positively chemotactic in certain concentrations became neutral or even negative when the concentration was altered.

We have seen that the action of the leukocytes in moving toward some substances and away from others is entirely analogous to similar phenomena occurring among lower, unicellular forms of life, and the explanations applied to the apparently conscious acts of the ameba, such as the motion toward and the engulfment of food, have been applied to the activities of the leukocytes as well. Many of the theories developed concerning the free living forms, however, have been easily excluded in the case of the leukocytes, because of the environment in which their activities are developed. Thus the many interesting reactions of paramecia and other organisms to light (heliotropism) have little bearing upon this subject, and the views based on the theories of orientation may be excluded on the ground of the symmetry of the normal leukocyte. The observations of Garrey,⁴¹ that indicate that it is the dissociated ions of various acids and bases which are responsible for the directive stimuli exerted upon certain flagellates, may yet result in throwing some light upon leukocytic movements, especially if we can come to accept the conceptions of ion-proteins upheld by Loeb⁴² and his pupils. However, the facts concerning these phenomena, as well as the possibility, previously mentioned, of the opposite electrical charges carried by the leukocytes and the substances attracting them cannot be regarded at present as more than interesting thoughts. Of more than merely speculative interest, however, are the views of chemotaxis which are based upon the study of conditions of surface tension. In order to consider these properly it will be useful to review briefly the fundamental principles governing these conditions.

The molecules of any fluid are held together by mutual attraction due to the force generally spoken of as cohesion. This force is exerted by like molecules upon each other in solids more strongly than in liquids, and in gases less strongly. Since we are dealing in this connection with occurrences taking place in liquids, we will restrict our consideration to these. The force of cohesion is influenced in a number of ways. Thus, for instance, heat reduces it, and this is the cause that solids are converted into liquids and liquids into gases, provided of course that the heat brings about no chemical change. In large masses of fluids the force of gravitation overcomes that of cohesion and larger masses of liquid assume the shape of the containing vessel. In smaller masses the force of cohesion tends to bring about the spherical shape. This comes about in the following way: Within the interior of a drop of liquid all the molecules attract each other, and since the force of attraction is equal in all directions it neutralizes itself, and the molecules are

⁴¹ Garrey. *Am. Jour. of Phys.*, 3, 1900.

⁴² Loeb. *Am. Jour. of Phys.*, 3, 1900.

uninfluenced by it, mobile and free. The molecules on the surface are in a different condition, however. They are subjected to the force of cohesion from within, but not from without, and are therefore drawn strongly toward the center. The result is the same as though the drop were subjected to pressure from without and the surface layers were in a state of compression. There is in consequence a constant tendency of all the surface molecules to be drawn toward the center and a resulting tendency to a diminution of the surface area. It is as though the surface of such a drop were a thin, elastic membrane which tended to contract and diminish in size and surface. The force with which this takes place is spoken of as surface tension,⁴³ and the energy underlying it is called, by Ostwald, surface energy. Since a drop of one fluid suspended in another with which it cannot mix is relieved of the disturbing factor of gravitation, its surface tension has the effect of contracting the small mass into a form which, for the given volume, will expose the smallest possible surface, and this is, of course, the sphere. It is for this reason that, if we shake up such systems as water and chloroform, or oil and water, the chloroform or the oil will be distributed through the water as small droplets. The degree of surface tension of any fluid is measurable by a number of reasonably accurate methods which may be found in any text-book of physics and which we need not consider here. It is of course dependent in each case upon the nature of the surrounding medium. We have taken into consideration above only the force which is exerted within the drop by the cohesion, that is, the attraction toward the center. This would be uninfluenced from without only in a vacuum. In nature the surface molecules, though forcibly drawn toward the center, are also affected from without by the attraction exerted by the molecules of the substances surrounding the drop. There is a constant balance, therefore, at any part of the surface of a drop of fluid between the cohesion tension from within and attractions from without. The resultant of the two forces determines the surface tension, which will be greater or less in inverse ratio to the attraction from without for any given drop, and a variation of the external attraction at different points on the periphery of the drop will naturally influence the shape of the drop. For a relief of attraction at one point would tend to permit that part of the surface to retract, and an increase in this attraction would tend to allow it to bulge, with the formation of a sort of pseudopod.

In studying the importance of surface tension⁴⁴ in determining the motions of unicellular organisms a number of important attempts have been made to imitate cell motion by means of the suspension of various substances of strong cohesive properties in liquid media. The

⁴³ Michaelis. "Dynamik der Oberflächen," Steinkopf, Dresden, 1909.

⁴⁴ For a thorough discussion of this phenomenon see also Gideon Wells, "Chemical Pathology," Saunders, 1911.

idea was suggested by Quinke,⁴⁵ and later by Bütschli,⁴⁶ but has been most extensively studied by Rhumbler.⁴⁷ The result has been the production of a number of "artificial amebæ" which in almost all respects behave like the living organisms. Thus if a small mass of mercury is placed into a dish filled with water acidified with nitric acid, and a small crystal of dichromate of potassium is dropped near the mercury, the dichromate will dissolve and a yellow cloud will gradually diffuse from it toward the mercury. As soon as the yellow cloud touches this it will begin to show change of form and to elongate in the direction of the dichromate, often moving toward it. The motion of the quicksilver will resemble with considerable accuracy that of an ameba moving toward a particle of food or sending out pseudopodia. A more striking and complete imitation is that obtained by Rhumbler when he placed a drop of clove oil into a mixture of alcohol and glycerin. The changes of surface tension produced upon the surface of the clove oil by the alcohol give rise to movements in the oil entirely analogous to those of motile cells in favorable media. The similarity has been extended even to the processes of engulfment of the food as observed among amebæ. Thus a drop of chloroform in water will flow about a particle of shellac and dissolve it. If a piece of glass coated with shellac is placed in contact with the drop it will engulf it, but will cast out the glass after the shellac coating has been dissolved away.

The similarity between phenomena purely referable to surface tension and those taking place in the living cells is therefore very striking and has been clearly analyzed in regard to its bearing upon leukocytic chemotaxis by Gideon Wells in his "Chemical Pathology." The chemotactic substances, diffusing to the leukocyte, will lower its surface tension on the side at which they come in contact. Pseudopodia will be thrown out on this side in consequence, and the leukocyte will move in this direction. The motion will be continued in this direction as long as the concentration of the chemotactic substance, and therefore the diminution of surface tension is greater on this side than on other parts of the periphery, until a point is reached at which the chemotactic substance is equally diffused on all sides, and motion will cease. The actual engulfment may then occur or the nature and concentration of the chemotactic substance may be so great that injury is done to the leukocyte. Whether or not the purely physical explanation of chemotaxis tells the whole story it is of course not possible to decide. At any rate, it furnishes a rational basis for

⁴⁵ Quinke. Quoted from H. G. Wells, "Chemical Pathology," Saunders, 1907.

⁴⁶ Bütschli. "Untersuch. über mikroskopische Schaume und das Protoplasma." Leipzig, 1892. See also H. G. Wells, *loc. cit.*, pp. 220 *et seq.*

⁴⁷ Rhumbler. *Arch. f. Entwicklungs Mechanik*, 1898.

the study of the phenomenon more promising than any of the others so far offered.

It is true, on the other hand, that such a theory in no way accounts for the apparently selective positive chemotaxis which is exerted by different substances. Thus the preponderance of polynuclear leukocytes in foci and serous cavities containing organisms like staphylococci, meningococci, streptococci, and others is in contrast to the lymphocytic accumulation in the pleural, subarachnoid, and peritoneal spaces infected with tubercle bacilli. Some writers have spoken, therefore, of active and passive leukocytosis according to whether or not the cells attracted seemed to possess ameboid motility. That surface tension phenomena alone do not account for this is clear. But it must be remembered that even tubercle bacilli, though eventually attracting few polynuclears and many lymphocytes, will cause an active polynuclear accumulation in the peritoneum and pleura when first injected, and are actively phagocyted. Later when the lesion is established and the bacilli are lodged in the tissues the polynuclears give way to the lymphocytes, which even then never accumulate in such proportion as do the microphages in acute suppurative lesions. It may well be that the chemotaxis originally attracting the polymorphonuclear leukocytes is the same in every case, but that a continued irritant, especially one well surrounded by tissue elements as are the organisms within the tubercles, may cease to exact any chemotactic influence, the accumulation of inactive lymphocytes possibly being due to a progressive death of these cells carried into the neighborhood of the lesion by the normal circulation of the lymph.

CHAPTER XII

THE RELATION OF THE LEUKOCYTES AND OF PHAGOCYTOSIS TO IMMUNITY

IN Metchnikoff's earliest work upon the daphnia or water flea he observed clearly that there was a direct relation between the degree of phagocytosis and the outcome of the infection. When phagocytosis of the invading yeasts was energetic and complete the daphnia recovered. When the yeast cells penetrated the intestinal wall of the daphnia in large numbers, and were enabled to multiply before the phagocytic cells could accumulate in sufficient numbers to engulf them, then the body of the daphnia was soon swamped with the parasites and death ensued.

This simple observation fostered the thought that the basic principle underlying all processes of immunity was represented in this struggle between the invading bacteria and the phagocytic cells. To the activity of the latter, entirely, he attributed the power of resistance.

In support of this contention Metchnikoff and his pupils have marshaled many facts, most of which are set forth in his classical work "*L'Immunité dans les Maladies Infectieuses*." It will be manifestly impossible here to do more than outline the plan of study which these investigations have followed and the conclusions to which they gave just foundation.

The original study upon the infectious disease of daphnia led to analogous experiments upon higher animals and, by the prolonged and patient investigations of Metchnikoff and his pupils, it was shown that, throughout the field of infectious disease, there was a striking parallelism between the resistance of the infected subject and the degree of phagocytosis which occurred.

Earlier studies concern themselves chiefly with the natural immunity possessed by many animals against certain infection. The infectious disease which at this time had been most thoroughly studied was anthrax, and Koch had shown that frogs and other cold-blooded animals were markedly resistant against this micro-organism. Taking advantage of this observation, Metchnikoff studied the phagocytosis of anthrax bacilli in frogs and found that it took place rapidly and effectively, all of the injected bacilli being soon engulfed by the accumulating cells. Similarly, active phagocytosis of anthrax

bacilli was demonstrated in such naturally resistant animals as dogs and chickens, while almost no cell ingestion occurred in delicately susceptible animals like guinea pigs and rabbits. Rats, on the other hand, more resistant to anthrax than guinea pigs, less so than dogs, showed a degree of phagocytosis intermediate between that observed in the cases of the other animals mentioned above. And yet, in these more susceptible animals, the normal bactericidal action of the blood upon anthrax bacilli, though never extreme, was often more marked than that of the naturally immune animals mentioned above.

It is well known, for instance, that the serum of dogs possesses almost no bactericidal properties for anthrax bacilli,¹ although the animals are highly resistant to this infection, while the serum of rabbits is probably more strongly bactericidal for these bacilli than the serum of most other animals, and yet rabbits are extremely susceptible. That the lack of bactericidal powers of the serum is not always a sign of susceptibility on the part of the animal was shown as early as 1889 by Lubarsch. (We must remember, however, that lack of bactericidal power does not necessarily mean lack of sensitizer. For bacteria may be sensitized without being killed extracellularly as can be shown by the alexin-fixation reaction.)

The study of anthrax infections was a peculiarly fortunate choice of subject, since in this bacillus resistance to serum lysis is especially well marked and phagocytosis seems indeed to be the chief mode of bacterial destruction. Studies analogous to those originally made with anthrax, however, were subsequently carried on with streptococci, pneumococci, and staphylococci chiefly by Bordet,² Marchand,³ and others, and results coinciding with those of Metchnikoff were obtained. In every case naturally resistant animals showed marked phagocytosis, and susceptible ones failed to show it to the same degree. It is a strong support of the same opinions, too, that Marchand's studies, later extensively confirmed, showed that the more virulent and invading strains of streptococci, the less active is the phagocytosis—a converse, but equally conclusive, observation.

Further support for this point of view is manifold and cannot be considered with anything like completeness. We may refer briefly, however, to the experiments of Vaillard, Vincent, and Rouget⁴ with tetanus, and those of Leclainche and Vallée⁵ with symptomatic anthrax, because they are especially valuable in illustrating the importance of phagocytosis in another class of infection. The poisons of these micro-organisms are extremely toxic for rabbits, and if

¹ Petterson. *Centralbl. f. Bakt.*, 1, 39, 1905.

² Bordet. *Ann. de l'Inst. Past.*, Vol. 11, 1897.

³ Marchand. *Archiv. de med. Exp.*, Vol. 10, 1898.

⁴ Vaillard, Vincent, and Rouget. *Ann. de l'Inst. Past.*, Vols. 5, 6, 1891-1892.

⁵ Leclainche and Vallée. *Ann. de l'Inst. Past.*, Vol. 14, 1900.

a small amount of culture material, together with agar, broth, or any foreign substance which may inhibit or divert phagocytosis from the spores, is injected into these animals rapid proliferation and death with toxemia result. If, on the other hand, the spores are carefully washed of foreign material and toxin rapid phagocytosis results and the animals recover.

The parallelism which was followed out so extensively between natural immunity and phagocytosis was even more closely marked in the case of artificially acquired immunity. The first observations of this kind made by Metchnikoff, again on the subject of anthrax infection, were carried out by the active immunization of rabbits. The subcutaneous injection of virulent anthrax bacilli into normal rabbits is usually followed by a rapid growth of the bacteria, with much serous exudation but hardly any leukocytic accumulation. In immunized animals, on the other hand, the bacilli are taken up by hosts of phagocytes, just as this occurs in naturally resistant dogs or other animals. Similarly Bordet⁶ has shown that cholera spirilla injected into the blood stream of cholera-immune animals are taken up by leukocytes even before they can be subjected to lysis by the circulating lytic antibodies.

It would add little to clearness were we to multiply the examples in which it has been demonstrated that the acquisition of increased resistance is accompanied by enhancement of the phagocytic process. This statement may be regarded as an axiom, and indeed our later discussions of the *opsonins* and *bacteriotropins* will show clearly why such a state of affairs is to be expected. Taken by itself, however, it does not necessarily prove that the destruction of the invading germs is entirely due to the leukocytes. It might still be possible that the bacteria are injured or even killed by the antibacterial serum constituents before they can be taken up and carried away by the cellular elements; the phagocytes then would act only as scavengers for the removal of the dead bodies. Indeed, this opinion was long held by a number of the adherents of the purely humoral school. However, such a point of view is no longer tenable—especially in the light of the later opsonin studies just referred to. Moreover, long before these more recent studies it was clear that bacteria may often grow within the leukocytes—finally destroying these—and that they may even remain fully virulent after ingestion. For, as Metchnikoff showed, if guinea pigs were injected with a little of the exudate formed after the injection of anthrax bacilli into immunized rabbits (an exudate in which there were no longer any extracellular bacteria because of energetic phagocytosis) death often resulted. It was clear, therefore, not only that the ingested bacteria were still alive, but that they were, at least in part, still fully virulent.

⁶ Bordet. *Ann. de l'Inst. Past.*, Vol. 9, 1895.

A further method of investigation employed by Metchnikoff in his endeavors to prove his point consisted in the attempt to demonstrate that virulent bacteria could be protected from destruction in the bodies of resistant animals if the leukocytes could be held at bay. This resulted in a number of ingenious experiments, the most convincing of which is the one carried out with anthrax bacilli and frogs by Trapetznikoff.⁷ Anthrax spores were inclosed in little sacks of filter paper and these were introduced subcutaneously into frogs. In consequence the spores, bathed in the tissue fluids, but protected from phagocytosis, developed into the vegetative forms, multiplied, and remained virulent for several days. Taken up by the frog's phagocytes under ordinary conditions, they would rapidly have been taken up, digested, and destroyed. Here again it was shown that the body of fluids alone were unable to dispose of the bacteria and that the natural resistance of the frog was due entirely to phagocytic processes.

Other experiments have been aimed at a general reduction of phagocytic activity by the use of narcotics. Thus, Cantacuzene⁸ showed that animals treated with opium are very much more susceptible to infection than are normal controls. And since opium markedly inhibits the activity of the white cells it may possibly be that these experiments furnish a further support for Metchnikoff's opinion. At any rate, it is worth noting that, even though these experiments are not convincing in their assertion that the increased susceptibility was due entirely to the interference with the leukocytes, they indicate very definitely the inadvisability of using morphin and similar narcotics in infectious diseases.

It is quite clear at any rate, then, that the process of phagocytosis increases in energy as immunity is acquired and, so far, Metchnikoff's assertions are entirely upheld by later knowledge. In his contention that *all* properties upon which the resistance of the animal against infection depends center directly or indirectly in the phagocyte, however, many subsequent amendments have been necessary, which will become self-evident in the following discussions of individual phases of the destruction of invading bacteria.

We cannot at the present time attempt to correlate these extreme views of Metchnikoff with the equally extreme opinions of those investigators who formerly attributed immunity entirely to the properties of the body fluids, assigning to the cellular activities a merely secondary rôle. Many of the apparently opposed contentions have become reconciled, and we now realize that neither process alone tells the whole story, both being parts of the complicated correlated processes which together constitute the mechanism of resistance. It was

⁷ Trapetznikoff. *Ann. de l'Inst. Past.*, 1891.

⁸ Cantacuzene. *Ann. de l'Inst. Past.*, Vol. 12, 1898.

indeed to the eager controversy between the two schools that we owe much of the clearness of conception which recent years have given.

After the bacteria are taken up by phagocytes they undergo a gradual disintegration or dissolution comparable to that by which a particle of food is digested within the cell body of a rhizopod. With the exception of such particularly insoluble micro-organisms as the tubercle bacillus, the leprosy bacillus, blastomyces, and a few others, there is in all cases an eventual complete resolution of the bacterial body. As in amebæ the digestion takes place often after the formation of digestive vacuoles, and by staining at this time with neutral red it may be demonstrated that the process goes on in a weakly acid environment.

Metchnikoff naturally assumed, therefore, that the intracellular digestion of bacteria by microphages (polynuclear leukocytes), or of cellular elements, etc., by macrophages, was a process carried on most probably by enzymes, and that these enzymes were identical with the bactericidal bodies described as "alexin" and "sensitizer" or "amboceptor" in the blood stream. To follow without confusion the development of his ideas, however, it is necessary to bear in mind that much of his earlier work was done at a time when the discovery of the coöperation of two substances in bacteriolysis and hemolysis (the amboceptor and the complement) had not yet been made by Bordet, and when the bactericidal effect of normal serum was attributed entirely to a single substance—the alexin of Buchner.

Buchner⁹ himself had suggested that alexin was an enzyme-like body probably derived from the leukocytes.

In his experiments Buchner had noticed that exudates, produced by intrapleural injections of aleuronat in rabbits and dogs, possessed a bactericidal value for *Bacillus coli* which exceeded the bactericidal power of the blood serum itself. The influence of active phagocytosis could be excluded by the fact that the leukocytes of the exudate had been killed by repeated freezing and thawing. Similar results were obtained by Hahn¹⁰ with *B. typhosus*.

Denys and Kaisin,¹¹ working along similar lines, found that the pleural exudates of rabbits, obtained by the injection of dead staphylococci and freed of cells by centrifugalization, were more highly bactericidal for staphylococci than the blood serum of the same animals, but found also that the inactivated exudate could not be reactivated by the addition of leukocytes. Denys offered as an explanation for these phenomena that the living leukocytes in the original exudate secreted alexin or complement which enhanced the bactericidal activity of the exudate, that the leukocytes, subsequently added to

⁹ Buchner. *Münch. med. Woch.*, No. 24, 1894.

¹⁰ Hahn. *Archiv f. Hyg.*, Vol. 25, 1895.

¹¹ Denys and Kaisin, Denys and Havet. *La Cellule*, Vol. 9, 1893; Vol. 10, 1894.

inactivated exudate, however, had lost vitality during the process of isolation and washing, and no longer possessed secretory power.

Hankin,¹² Kanthack and Hardy¹³ had gone even farther than this, and had attributed the production of alexin to the eosinophile leukocytes particularly.

Metchnikoff,¹⁴ basing his opinion on his own studies, those of his pupils, and many other investigations similar to those mentioned above, came to the conclusion that, under ordinary conditions, the normal blood contains no free bactericidal substances. He assumes that these substances are entirely intracellular, being constituents of the various phagocytic elements, by means of which the cells digest the foreign elements they take up. He believes that there are essentially two varieties of such digestive enzymes or "cytases"—just as there are two varieties of phagocytes. The microphages, chiefly concerned in the digestion of bacteria, secrete the bactericidal alexin, or, as Metchnikoff calls it, "microcytase." The macrophages, the large mononuclear lymph and endothelial cells, primarily concerned in the phagocytosis of cellular elements (red cells, etc.), contain another variety of digestive enzyme, the "macrocytase," or cytolytic (hemolytic) alexin. The supposition that the hemolytic "cytase" is largely derived from the macrophages was based particularly upon the investigations of Metchnikoff's pupil, Tarassewitch,¹⁵ who found that the extracts obtained from lymph nodes, and other organs rich in macrophages, possessed hemolytic properties. Both this work and the preceding studies regarding the extraction of alexin from polynuclear leukocytes will be more fully discussed below.

Maintaining that these cytases are purely intracellular under ordinary conditions, Metchnikoff believes that, in normal animals, the destruction of invading bacteria or of injected cellular substances (blood cells, etc.) is accomplished entirely by the phagocytic process, with subsequent intracellular digestion. In immunized animals, however, there is present in the circulating blood another substance, not identical with the cytases, but also derived from the leukocytes or from the blood-forming organs—the "fixateur" (Ehrlich's "amboceptor"—Bordet's "sensitizer"). This specific "fixateur" sensitizes the bacteria or other antigens to the action of the cytases. For his assumption regarding the origin of this sensitizer he finds support largely in the researches of Pfeiffer and Marx, and others mentioned in our section on the origin of antibodies, as well as in the similar investigations of Deutsch,¹⁶ carried on under Metchnikoff's personal supervision.

¹² Hankin. *Centralbl. f. Bakt.*, Vol. 12, 1892.

¹³ Kanthack and Hardy. *Proc. Roy. Soc.*, Vol. 52, 1892.

¹⁴ Metchnikoff. *Ann. de l'Inst. Pasteur*, Vol. 7, 1893; Vol. 8, 1894; Vol. 9, 1895.

¹⁵ Tarassewitch. *Ann. de l'Inst. Past.*, Vol. 16, 1902.

¹⁶ Deutsch. *Ann. de l'Inst. Pasteur*, Vol. 13, 1899.

Final digestion of the sensitized antigens (bacteria or blood cells), however, can take place only under the influence of the cytases intracellularly, unless by previous leukocytic injury these enzymes have been liberated into the blood stream.

While it is admitted, then, that bacteria may be killed and digested both intra- and extracellularly in the animal body, the cytases, which accomplish this, are regarded as the same in both cases, being derived from the phagocytic cells. In immunized animals "fixateur" may be produced under the stress of active immunization and furnished to the blood stream by the blood-forming organs. By this substance bacteria and cells may be sensitized. However, the enzyme by which digestion is actually accomplished, "cytase" or alexin, is not present free in the blood even in immune animals unless it has become free and extracellular by injury to the leukocytes.

How, then, on this basis does Metchnikoff account for the Pfeiffer phenomenon, in which the extracellular destruction of bacteria takes place rapidly in the peritoneal exudate? His explanation is the following: When bacteria or other substances are injected into the peritoneum there is a preliminary injury of leukocytes (phagolysis), and by this alexin or cytase is liberated. When cholera spirilla, for instance, are injected into the peritoneal cavity of an immunized guinea pig there follows a short period during which few if any leukocytes are present in the exudate, but many may be found gathered in motionless clumps in the folds of the peritoneum and mesentery, incapable of phagocytosis and apparently injured. If such leukocytic injury can be avoided Metchnikoff claims that the extracellular lysis of cholera spirilla will fail to take place. Thus if sterile broth or salt solution is injected into the peritoneum of a guinea pig a preliminary phagolysis will be followed by an accumulation of leukocytes. If cholera spirilla are now introduced no extracellular digestion is seen, but, instead of this, rapid phagocytosis takes place. This he says is due to the fact that the freshly accumulated, healthy phagocytes, collected in response to the preliminary broth injection, are not easily injured and do not undergo phagolysis; no cytase is liberated and, in consequence, no serum bacteriolysis can take place. In the same way he claims that the hemolysis of red blood cells (goose blood) in the peritoneum of specifically immunized guinea pigs may be prevented if, by a previous injection of broth, healthy leukocytes have been caused to accumulate. In such a case the goose blood corpuscles are rapidly ingested by the phagocytes and no hemolysis occurs.

It is self-evident that the validity of this interpretation of the occurrences is strictly dependent upon the demonstration that the circulating blood normally contains no alexin or complement. This is rigidly maintained by Metchnikoff, and is indeed one of the most important uncertainties of serology. He asserts that alexin appears

in the blood serum only because changes in the leukocytes take place during coagulation. It is not, by any means, settled that Metchnikoff is right in this—in fact, more recent investigations seem to show that he is wrong, and that we may assume definitely that alexin is present in considerable amounts in the circulating blood plasma of normal animals.

Metchnikoff's denial of this is based chiefly on the experiments of Gengou. Gengou¹⁷ took the blood from various animals into paraffined tubes and centrifugalized it at low temperature before it could clot. This freed the plasma from the cells before clotting, though coagulation of course took place as soon as this plasma was removed to tubes and kept at room temperature. The serum expressed from this clotted plasma he compared for alexin contents (bactericidal properties) with that obtained from clotted whole blood.

He found that, in all cases examined (dogs, rabbits, and rats), the plasma contained practically no bactericidal substances, or at any rate an incomparably smaller amount than was present in the serum obtained from the clotted blood.

These experiments of Gengou would be conclusive in establishing Metchnikoff's theory if they were confirmed by other observers. This, however, has not been the case. Petterson¹⁸ found no difference between the bactericidal properties of serum and oxalate plasma, and Lambotte¹⁹ arrived at similar results when he compared serum with the coagulable plasma obtained by tying off a section of a vein and centrifugalizing the blood without opening the vessel. Hewlett,²⁰ Falloise,²¹ Schneider,²² and more recently Dick²³ and Addis,²⁴ whose work has been done with particular attention to technical accuracy, fail to confirm Gengou, finding no appreciable difference between plasma and serum.

In favor of Gengou's results are the investigations of Herman²⁵ and the more recent ones of Gurd.²⁶ Further supporting Gengou's conclusion is the observation recorded by a number of workers (Walker,²⁷ Longcope,²⁸ and others) that the complement or alexin contents of serum will increase somewhat as the serum is allowed to

¹⁷ Gengou. *Ann. de l'Inst. Past.*, Vol. 15, 1901.

¹⁸ Petterson. *Arch. f. Hyg.*, Vol. 43, 1902.

¹⁹ Lambotte. *Centralbl. f. Bakt.*, Vol. 34, 1903.

²⁰ Hewlett. *Zeitschr. f. Heilkunde*, 24, 1903.

²¹ Falloise. *Bull. de l'Acad. Roy. de Méd.*, 1905.

²² Schneider. *Arch. f. Hyg.*, 65, 1908.

²³ Dick. *Jour. Inf. Dis.*, Vol. 12, 1913.

²⁴ Addis. *Jour. Inf. Dis.*, Vol. 10, 1912.

²⁵ Herman. *Bull. de l'Acad. Roy. de Méd.*, 1904.

²⁶ Gurd. *Jour. Inf. Dis.*, Vol. 11, 1912.

²⁷ Walker. *Jour. of Hyg.*, 3, 1903.

²⁸ Longcope. *Med. Bull. Univ. of Pa.*, 1902, Vol. 15, p. 331.

stand on the clot. This observation, too, has been rendered inconclusive by contrary reports from other investigators. Longcope,²⁹ further, has found that alexin was more plentiful in the blood of individuals suffering from leukemia—in which of course a larger percentage of leukocytes is present in the circulation. This, too, has been contradicted by other workers, but even if upheld would not influence the possibility of there being alexin in the normal circulation. On the whole Gengou's contentions with their consequent bearing upon Metchnikoff's theory cannot be accepted as final. In fact, the greater part of available experimental evidence seems to point to the actual presence of alexin in the normal circulating blood. This seems also indicated by the unquestionable fact that active phagocytosis may take place in the circulation of an animal and, as we shall see below, free alexin is probably necessary (as opsonin) in this process. Further evidence in this direction also is furnished by the immediate anaphylactic shock which follows the injection of antigen into the blood stream of a sensitized animal, a process in which we have much reason to believe that alexin takes an active part. However, the problem is a difficult one, and, while we favor the opinion that free alexin is present in the intravascular blood, we must admit that a crucial experiment has not yet been formulated.

Now, as regards the apparent extraction of alexin from leukocytes and lymphatic organs, we have already called attention to the fact that most of the researches associating these cells with the bactericidal substances were carried out before the dual mechanism of sensitizer and alexin in bacteriolysis had been fully worked out. In consequence conclusions were formulated from the mere facts of the presence of bactericidal or hemolytic properties in cell-extracts without the further determination of heat stability or the possibility of reactivation. Most of the earlier work also was done without sufficient attention to the separation of the cells and the serum of leukocytic exudates. The first one to do this carefully was Hahn,³⁰ who, like his predecessors, concluded that the bactericidal leukocytic substances, undoubtedly encountered by him, were identical with alexin. Doubt was first cast upon this by Schattenfroh,³¹ who worked with leukocytes suspended and extracted in physiological salt solution. He found that bactericidal substances were, indeed, obtained in this way, but that, unlike alexin, these substances were thermostable, withstanding exposure to a temperature of 56° C. and destroyed only by exposure to temperatures as high as 75° C. to 80° C. for thirty minutes.

²⁹ Longcope. *Jour. of Hyg.*, Vol. 3.

³⁰ Hahn. *Arch. f. Hyg.*, Vol. 25.

³¹ Schattenfroh. *Arch. f. Hyg.*, Vols. 31 and 35, 1897.

Moxter,³² a little later, working with cholera spirilla, also came to the conclusion that the leukocytic bactericidal substances were not identical with those found in the blood serum. Petterson,³³ too, made thorough investigations into the nature of the bactericidal substances extracted from the leukocytes, and, working chiefly with *B. proteus* and *B. anthracis*, found such substances in the leukocytes of dogs, rabbits, and guinea pigs active against the bacteria mentioned above, but failed to find them active, at least in guinea pig and cat leukocytes, against *B. typhosus* or the cholera spirillum. He expresses the opinion that bactericidal leukocytic substances are normally given up to the blood in minute quantity only or not at all, and that these substances hold no definite relationship to the bactericidal substances found in blood serum. In a later investigation he showed that the "endolysins," as he now calls the leukocytic bactericidal substances, may, like many enzymes and serum bacteriolysins, be precipitated out of solution with alcohol and ether; but he separates them absolutely from serum lysins and complement. The latter, while they may be, in part at least, secreted by the leukocytes, are, according to Petterson, induced easily to come out of the cells during life by slight injury or other stimulation, while the endolysins themselves are abstracted from the cells only after extensive extraction or maceration.

Schneider³⁴ has come to similar conclusions and speaks of the endocellular bactericidal substances as "Leukine." In a recent investigation of the same subject the writer³⁵ has in all essentials confirmed Schattenfroh's original conclusions regarding the heat stability of the extracted leukocytic bactericidal substances, and has shown that after inactivation by heat these substances are not reactivable by the addition of fresh leukocytic extracts, and that the yield obtained from the leukocytes of immunized animals is not greater than that obtained from normal leukocytes.

It appears, therefore, that, contrary to Metchnikoff's first supposition, the enzymes which bring about the digestion of phagocytized bacteria within the cell are not identical with those which bring about a similar extracellular digestion. In addition to the demonstration of a definitely different structure possessed by the bactericidal leukocytic extracts, as evidenced by their heat stability, we have the negative evidence that neither true alexin nor sensitizers have ever been successfully extracted from such cells.

It is still possible that this may eventually be done—but, although indirect evidence like that of Denys, Longcope's observations in leukemia, and the occasional increase of the alexic powers of serum

³² Moxter. *Deutsche med. Woch.*, 1899, p. 687.

³³ Petterson. *Centralbl. f. Bakt.*, i, 39, 1905; 46, 1908.

³⁴ Schneider. *Archiv f. Hyg.*, Vol. 70, 1909.

³⁵ Zinsser. *Jour. Med. Res.*, Vol. 22, 1910.

after standing on the clot points to a probability of this, no direct evidence has so far been satisfactorily produced. In the hope that the leukocytes would give up alexin—possibly as a secretion as suggested by Denys—the writer, with Cary, some years ago kept washed leukocytes at 37.5° C. in Locke's solution, but was unable to find any evidence of alexin production within 48 hours.

The apparent extraction of hemolysin from macrophages by Tarassewith, moreover, has met with a similar refutation. Korschun and Morgenroth³⁶ have shown that these hemolytic extracts are extremely heat resistant, are alcohol and ether soluble, and do not act as antigens. They are quite different from the serum hemolysins, therefore, and probably closely related to the hemolytic lipoidal substances described by Noguchi and others.

Further strong arguments against the assumption of the presence of hemolytic alexin in the body of the macrophages have been advanced by Gruber³⁷ and by Neufeld.³⁸ Gruber showed that no extracellular hemolysis takes place when leukocytes are brought together with sensitized red blood cells, and Neufeld showed that even after the phagocytosis of such sensitized cells the hemolysis is very much slower, and of a different character from extracellular serum hemolysis. In the intracellular digestion there are no mere solution of the hemoglobin and formation of a shadow form (stroma), but there occur a gradual degeneration with the formation of a granular detritus of hemoglobin.

It is probable, then, that the digestion of bacteria and cells within the phagocytes is carried out by substances not identical with those taking part in serum lysis. It is not unlikely that the intracellular process is a quite complicated one, not depending on a single enzyme.

In addition to the bactericidal substances extracted from leukocytes a number of true enzymes have indeed been obtained by various investigators. We have mentioned in another place that one of the earliest observations in this respect was that of Leber,³⁹ who noticed that sterile pus could liquefy gelatin. It may be commonly observed in paraffin or celloidin sections of staphylococcus abscesses that a ring of apparently digested or degenerating tissue is formed about an accumulation of leukocytes—in foci in which the bacteria may be too sparse to be held accountable for the changes. These leuko-proteases have subsequently been carefully studied by Opie,⁴⁰ Jochmann and Müller,⁴¹ and a number of others.

³⁶ Korschun and Morgenroth. *Berl. klin. Woch.*, 1902.

³⁷ Gruber. Quoted from Sachs, in "Kraus u. Levaditi Handbuch," Vol. 2, p. 991.

³⁸ Neufeld. *Arb. a. d. kais. Gesundheits. Amt.*, Vol. 28, 1908.

³⁹ Leber. "Die Entstehung der Entzündung," Leipzig, 1891.

⁴⁰ Opie. *Jour. Exp. Med.*, Vol. 7, 1905; Vol. 8, 1906; Vol. 9, 1907.

⁴¹ Müller and Jochmann. *Münch. med. Woch.*, Nos. 29 and 31, 1906.

Opie found that two distinct proteolytic enzymes could be extracted from the cells of exudates obtained by turpentine injections. One—peculiar to the polynuclear leukocyte, and similar to one previously described by Müller ⁴²—acts in a weakly alkaline medium. The other, present particularly in exudates containing a predominating number of mononuclear cells, acts in a weakly acid reaction. Tschernorutski also found proteolytic ferments in both micro- and macrophages, but found no lipase in the polynuclear extracts. This seems particularly interesting in view of the great resistance to intracellular digestion noticed in acid-fast bacteria, a point of some importance in connection with the destruction in the body of such micro-organisms as the bacilli of tuberculosis and leprosy.⁴³ Jochmann ⁴⁴ states that the action of the leukoprotease, which acts in an alkaline medium upon casein, results in the formation of tryptophan and ammonia, and believes it to be functionally very similar to trypsin. It is interesting to note that the most active protease is obtained from pus as it forms about acute infections or other stimuli which lead to the accumulation of polynuclear leukocytes, whereas it is apparently completely absent from tuberculous pus.

The question immediately arises, are these leukoproteases identical with the bactericidal substances extracted from leukocytes as described above? For it might well be that bacterial death resulted merely from the digestive action of the enzyme upon the bacterial protein. Jochmann,⁴⁵ who has approached this problem experimentally, has answered it in the negative. By repeated alcohol precipitation of glycerin extracts of leukocytes he obtained an enzyme preparation which possessed absolutely no bactericidal properties, though it was still actively proteolytic.

Not only did this relatively pure ferment possess no bactericidal action, but bacteria actively proliferated when suspended in it. Jochmann believes that living bacteria are not amenable to the enzyme possibly because of their possession of an "antiferment," at least this would follow in some cases from the experiments of Kantorowicz.⁴⁶

The leukoproteases, therefore, appear to possess no direct significance in bacterial immunity. Their function seems rather to lie in the resorption of dead tissues, fibrin, blood clots, etc. Friedrich Müller ⁴⁷ has pointed out their possible importance in the rapid destruction and liquefaction of the massive fibrinous exudates remaining after the crisis in lobar pneumonia.

⁴² Müller. *Congr. f. inn. Med.*, Wiesbaden, 1902.

⁴³ Zinsser and Cary. *Jour. A. M. A.*, 1911.

⁴⁴ Jochmann. *Leucozyten Fermente*, etc., "Kolle u. Wassermann Handbuch," 2d Ed., Vol. 49, 2.

⁴⁵ Jochmann. *Zeitschr. f. Hyg.*, 61, 1908.

⁴⁶ Kantorowicz. *Münch. med. Woch.*, No. 28, 1909.

⁴⁷ Friedrich Müller. "Verhand. d. Kongr. f. inn. Med.," 1902.

From the discovery of antibacterial properties in the extracts of leukocytes it is but a logical step to the attempt to utilize these substances therapeutically. This was especially called for in view of the disappointing results which have attended the injection of even large amounts of bactericidal sera into animals and human beings in whom anthrax bacilli, streptococci, or any other of the invasive bacteria or true parasites had gained a foothold. Petterson⁴⁸ was probably the first to study this phase of the problem systematically in connection with anthrax infection in dogs and rabbits. In preliminary studies he claimed to have determined that when leukocytes are left in contact with serum for four hours or longer there develops in the mixture a bactericidal power far superior to that which is possessed by these elements when separately kept in salt solution and mixed only just before the bactericidal tests. He attributes this to the fact that in dogs, at least, the leukocytes furnish bactericidal substances to the serum—an assumption which is entirely in accord with the earlier opinion of Denys and Kaisin,⁴⁹ which we have mentioned in another place. In direct continuance of these experiments he injected leukocytes into dogs at the same time at which he infected them with anthrax and observed a moderately protective influence, which, however, he admits was not very great. He followed this work in 1906 with similar observations on the protective influence of leukocytes in intraperitoneal infections of guinea pigs with typhoid bacilli. In these experiments⁵⁰ he made the curious observation that, although such protective influence was unquestionable, the guinea pig leukocytes contained no bactericidal substances active against typhoid bacilli. In consequence he concluded that the destruction of these bacteria in the guinea pig was due entirely to the serum-antibodies absorbed by the micro-organisms before phagocytosis, even when the actual destructive process took place intracellularly. The protective effect following on the injection of the leukocytes he attributed to an indirect influence of the leukocytic substances in stimulating the more rapid accumulation of alexin or complement in the peritoneum, with consequently more powerful phagocytosis. Following this, in 1908, Opie⁵¹ carried out experiments in which he observed that leukocytes injected intrapleurally into dogs, together with tubercle bacilli, exerted a distinct protection in that the course of the disease was prolonged and the tendency toward healing was more pronounced than in the controls.

In the same year extensive observations on the protective properties of leukocyte extracts were published by Hiss.

⁴⁸ Petterson. *Centralbl. f. Bakt.*, Vol. 36, 1904.

⁴⁹ Denys and Kaisin. "La Cellule," Vol. 9, 1893.

⁵⁰ Petterson. *Centralbl. f. Bakt.*, Vols. 40 and 42, 1906.

⁵¹ Opie. *Jour. Exp. Med.*, 1908.

Hiss⁵² worked at first with extracts of dog, rabbit, and guinea pig leukocytes; later he confined himself entirely to rabbit leukocytes. He extracted the leukocytes at first by repeated freezing and thawing in physiological salt solution, but the technique of his subsequent work was uniformly as follows: Intrapleural injections of aleuronat emulsions were made in rabbits and, after about 24 hours, the resulting exudates were taken away with sterile pipettes and centrifugalized before clotting could take place; the serum was decanted and the leukocytes then emulsified in distilled water, in quantity about equal to the amount of serum poured off. In this the leukocytes were allowed to stand for a few hours at incubator temperature, and then in the ice-box until used. For his experimental work in both animals and man, in most instances, not only the clear supernatant fluid was injected, but the cell residue as well; since Hiss realized that the extractions were necessarily incomplete. In intravenous work, of course, the supernatant fluid alone was injected.

With leukocytic extracts so prepared Hiss treated staphylococcus, typhoid bacillus, pneumococcus, streptococcus, and meningococcus infections in rabbits and obtained results which justified him in concluding that the leukocyte extract exerted strong protective action in all of these cases. Many of his animals survived infections fatal to controls even when the treatment was delayed as long as 24 hours after infection. Subsequently Hiss and Zinsser⁵³ treated series of patients, ill with pneumonia, meningitis, and staphylococcus infections, with leukocyte extracts prepared by the method of Hiss, and felt that they were justified in concluding that in many cases, at least, the course of the disease was favorably influenced by the leukocyte extract. Favorable results have since then been obtained also by Lambert in erysipelas, and by Hiss and Dwyer in a variety of conditions. Dwyer has used the extract in various infections of the eye, ear, nose, and throat.

While there seems to be little question about the actually favorable influence of the leukocyte extract, both in experiments with animals and in the treatment of human cases, there has been considerable difficulty in determining the reasons for this influence. In subsequent studies Hiss and Zinsser (*loc. cit.*) were able to show that the extracts did not favor phagocytosis and that the moderate bactericidal properties possessed by the leukocytic substances could not account for their effectiveness. There *did* seem to be a more rapid accumulation of phagocytes in the peritoneal cavities of guinea pigs infected with cholera spirilla when leukocyte extract was injected with the bacteria, and it is not impossible, in fact, it seems probable to the writer, from subsequent experience, that the protective prop-

⁵² Hiss. *Jour. of Med. Res.*, new series, Vol. 14, 1908.

⁵³ Hiss and Zinsser. *Jour. of Med. Res.*, new series, Vol. 14, 1908.

erties of the leukocyte extracts are attributable, in part at least, to their positively chemotactic effect.

The entire problem opened up by the work of Hiss cannot be regarded as settled. Observations, both experimental and clinical, are still in progress, and it is hoped that the next few years may definitely decide in how far this treatment is applicable to human cases. It is not easy to draw conclusions from clinical observations since it is impossible to parallel such cases with untreated controls; in consequence the truth can be elucidated only by a multiplication of statistics. While the writer and others have treated a great many cases with disappointment, again the striking results occasionally observed have been so encouraging that it seems of the utmost importance to give the treatment extensive trial, especially since many injections have been made without any harm whatever to the patients. Although the experience thus far gathered permits of no definite conclusions, the writer would suggest from his own experience and his observation of that of others that the use of the leukocyte extract of Hiss be confined for the present to diseases like erysipelas, meningitis, and the pyogenic infections in which the process is distinctly localized and no general septicemia has supervened. It should also be given a thorough trial in broncho- and lobar pneumonia in which the bacteriemia which occurs represents very probably a constant discharge into the blood stream of bacteria from the pneumonic focus, rather than the firm establishment of bacterial growth within the blood itself. With few exceptions absolutely no results seem to have followed its use when such a septicemia has become established. In the class of cases first mentioned, however, where a localized infection has been obstinate and unusually violent, many brilliant results have been obtained. Judging from the results of Dr. Adrian Lambert, and more recent ones obtained by Dr. Dwyer, we would have no hesitation in stating that erysipelas is favorably influenced in most cases. The above suggestions are made since it seems that in the question of clinical therapy much delay in the proper estimation of the value of a new type of treatment can be avoided by an intelligent choice of cases.

CHAPTER XIII

FACTORS DETERMINING PHAGOCYTOSIS

OPSONINS, TROPINS

FROM the very beginnings of his researches upon phagocytosis Metchnikoff recognized that the process was profoundly influenced by the properties of the fluid constituents of the blood plasma in which the phenomenon occurred. Both he and his pupil Bordet,¹ at this time working in Metchnikoff's laboratory, noticed that the phagocytic activity of leukocytes was greater in immune than in normal sera and associated this with the specific properties of the immune substances or antibodies in these sera; Metchnikoff himself interpreted the phagocytosis-enhancing power of the serum as a stimulation of the leukocytes and referred to the serum constituents by which this effect was produced as "stimulins." A closer analysis of the factors involved in this interrelationship, however, was not attempted at this time by him or his pupils, although indirect reference was made to it in a number of articles emanating from this school in the course of investigations on kindred problems of phagocytosis. Thus Gabritschewsky,² in 1894, published a paper on "Leukocytose dans la Diphthérie," in which he concluded that the poison of diphtheria bacilli, among other harmful effects, diminished the phagocytic power of the leukocytes, and that one of the beneficial influences of the curative serum was to render these and other cells "less sensitive to the bacterial poisons." This may be interpreted as indicating an assumption that the action of an immune serum in increasing phagocytic activity rested rather upon its influence upon the bacterial products than upon any stimulation of the phagocytes themselves. However, in diphtheria the action of the leukocytes was, even at this time, recognized as a merely secondary one, and Gabritschewsky's results did not materially influence the "stimulin" conception.

The first extensive investigation which occupied itself directly with these problems was that of the Belgian bacteriologists Denys and Leclef.³ The publication of these workers deals primarily with

¹ Bordet. *Ann. de l'Inst. Past.*, 1895.

² Gabritschewsky. *Ann. de l'Inst. Past.*, 1894.

³ Denys and Leclef. *La Cellule*, 11, 1895.

the nature of streptococcus immunity in rabbits. It established, first of all, the paramount importance of phagocytosis in the resistance of animals against these bacteria, and made clear that the destruction of bacteria was carried out equally as well by the leukocytes of normal as by those of immune animals, but was powerfully enhanced when either normal or "immune" leukocytes were combined with immune serum. Their work, therefore, indicated again that the increased phagocytosis of virulent bacteria, taking place in immune animals, depended clearly upon alterations in the functions of the serum rather than in those of the cells, and they suggested that the influence of this serum was not necessarily one of leukocyte stimulation, but might rather consist in action upon the bacteria, rendering them less resistant to phagocytosis. They say in substance: "À notre avis, on pourrait tout aussi bien admettre que la substance vaccinante ou antitoxique agit, non pas sur le leukocyte, mais sur un poison renfermé dans le corps du microbe ou dissous dans le milieu, et qui préserve le micro-organisme contre les atteintes du leukocyte."⁴

In this statement we have, in brief, the distinct formulation of our present view of the "opsonins."⁵

Observations with pneumococci and streptococci carried out after this by Marchand⁶ and by Mennes,⁷ whose investigations we cannot discuss in detail, beside confirming most of the observations of Denys and Leclef, brought out especially the relation of the virulence of micro-organisms to phagocytosis, showing that very virulent strains were taken up to a slight degree only in the presence of normal serum, but were subject to active phagocytosis when immune serum was employed. This, too, seemed to point primarily to the fact that the serum influenced rather the bacteria than the phagocytes, although no convincing proof is brought for this in their publications. Though much that had bearing indirectly on this problem was written during the following years, no definite progress was made beyond the results of Denys and his pupils until 1902,

⁴ In our opinion one can just as well believe that the vaccinating or anti-toxic substance acts not upon the leukocyte but upon a poison inclosed within the body of the bacteria or dissolved in the medium, which preserves the micro-organism against the attacks of the leukocyte.

⁵ Denys formulated this view with still greater clearness and positiveness at the Congress of Hygiene held at Brussels in 1903. We take our citation from the discussion on opsonins by Gruber (3d meeting Freie Vereinigung f. Mikrobiol., Vienna, 1909, *Centralbl. f. Bakt.*, I Ref., Vol. 44, Suppl. p. 3). Following is Denys' statement: 1. The phagocytosis in immune sera is dependent upon substances which are precipitated with the euglobulins. 2. These substances cause phagocytosis by inciting a physical alteration of the micro-organisms. 3. These substances are specific.

⁶ Marchand. *Arch. de Méd. Exp.*, 1898.

⁷ Mennes. *Zeitschr. f. Hyg.*, Vol. 25.

when Leischman⁸ introduced a technique by means of which it became possible to observe the process of phagocytosis with fresh serum and leukocytes *in vitro*.

By utilizing this technique and improving upon it Wright and Douglas in the following year (1903) evolved a method by means of which phagocytic activity could be quantitatively measured with reasonable accuracy. They worked at first with staphylococcus phagocytosis by human leukocytes in the presence of human citrate plasma, a research undertaken primarily because Wright,⁹ in collaboration with Windsor, had previously determined that human blood serum possessed practically no bactericidal power for this organism, and that phagocytosis was probably the chief mechanism of protection which the human body possessed against these bacteria. The researches of Wright and Douglas¹⁰ were carried out chiefly by mixing equal volumes of bacteria, serum, and leukocytes (in citrate suspension),¹¹ allowing these elements to remain together at 37.5° C. for varying periods, then staining on slides and determining the degree of phagocytosis by counting the numbers of bacteria taken up by each polynuclear leukocyte. Though many technical difficulties had to be overcome, and although the method at its best still permits of much personal error, careful work and untiring repetition made possible a considerable degree of accuracy, and definite facts regarding the mechanism of phagocytosis, heretofore merely suspected, could be recorded. The most important result of these investigations was the unquestionable establishment of the function of serum in the process of phagocytosis, namely, that it in no way "stimulated" the leukocytes in the sense of Metchnikoff, but rather acted entirely upon the bacteria, preparing them for ingestion. For this reason Wright coined the word "opsonins" (*ὀψονέω* = I prepare food) for the serum constituents which brought about this effect, believing them to be new antibodies, entirely distinct from the other serum antibodies heretofore recognized.

Wright and his followers now concluded that the rôle of the leukocyte in taking up bacteria was entirely dependent upon the opsonin contents of the serum. In a menstruum containing no serum, or in a serum in which the opsonins had been destroyed by heat, they found practically no phagocytic action on the part of washed serum-free leukocytes, and they, therefore, doubted the occurrence of spontaneous phagocytosis on the part of leukocytes themselves.

⁸ Leischman. *Brit. Med. Jour.*, Vol. 2, 1901, and Vol. 1, 1902.

⁹ Wright and Windsor. *Jour. of Hyg.*, Vol. 2, 1902.

¹⁰ Wright and Douglas. *Proc. Roy. Soc.*, 72, 1903, 73 and 74, 1904. See also Wright, "Studien über Immunisierung," Fischer, Jena, 1909.

¹¹ At first bacteria were merely mixed in equal volumes with citrated whole blood.

In this point it is not unlikely that Wright is mistaken, since other observers, notably Löhlein,¹² have observed the phagocytosis of various bacteria by washed leukocytes in indifferent, opsonin-free media. Although we may take it as assured that such spontaneous phagocytosis may take place (Metchnikoff and a number of others having obtained results similar to those of Löhlein), this is probably never very intense.

In fact, Wright, in some of his later work, does not insist rigidly upon the non-occurrence of spontaneous phagocytosis, but attempts to associate such phenomena with the salt contents of the medium in which it occurs. Together with Reid,¹³ he determined that spontaneous phagocytosis of tubercle bacilli unquestionably takes place, is most intense at a concentration of about 0.6 per cent. NaCl, and diminishes as the concentration is increased. This, as we shall see, has bearing on the possible physical explanations advanced to account for opsonic action, and has its parallels in experiments on the influence of electrolytes on agglutination and precipitation.

The fact remains that Wright demonstrated by his work that Metchnikoff's original view, which interpreted the difference between susceptibility and immunity as a difference between the inherent phagocytic powers of the leukocytes, is incorrect, and that the essential regulating influence affecting phagocytosis rests upon the action of the serum upon the bacteria.

The following experiment from the work of Hektoen and Ruediger¹⁴ illustrates this point with exceptional clearness. It shows that human leukocytes in the presence of normal defibrinated blood will take up bacteria energetically. When the leukocytes, however, are washed free of blood and added to untreated bacteria phagocytosis is practically nil. If, however, such washed leukocytes are mixed with bacteria that have been previously in contact with serum active phagocytosis will take place. In other words, the bacteria have been altered by the serum in such a way that they are now amenable to phagocytosis by washed leukocytes. The serum then acts upon the bacteria and not upon the leukocytes.

TABLE II
Phagocytosis by Human Leukocytes of Sensitized Bacteria

| | Average Phagocytosis |
|--|-------------------------|
| Human leukocytes (defibrinated blood) + <i>Staphylococcus aureus</i> | 22. |
| Human leukocytes (washed in NaCl solution) + <i>Staphylococcus aureus</i> | 1.2 |
| Human leukocytes (washed in NaCl solution) + <i>Staphylococcus aureus</i> (treated with human serum)..... | 10. |
| Human leukocytes (defibrinated blood) + <i>Streptococcus 300</i> | 22. |

¹² Löhlein. *Centralbl. f. Bakt.*, 38, 1906, Beiheft, p. 32; also *Münch. med. Woch.*, 1907, p. 1473.

¹³ Wright and Reid. *Proc. of Royal Soc. B.*, Vol. 77, 1906.

¹⁴ Hektoen and Ruediger. *Jour. Inf. Dis.*, Vol. 2, 1905, p. 132.

| | Average Phagocytosis |
|--|-------------------------|
| Human leukocytes (washed in NaCl solution) + Streptococcus 300 . . . | 1. |
| Human leukocytes (washed in NaCl solution) + Streptococcus (treated with human serum) | 14. |
| Human leukocytes (washed in NaCl solution) + Streptococcus (treated with guinea pig serum) | 12. |
| Human leukocytes (washed in NaCl solution) + Streptococcus (treated with rabbit serum) | 14. |

Wright and Douglas'¹⁵ work was done at first with normal serum or normal citrate plasma, and in this case they found that the opsonins were essentially unstable, being easily weakened by exposure to light, or heat, and even when preserved in sealed tubes in the dark they diminished noticeably on standing for 5 or 6 days. Other writers who have worked with the opsonic substances in normal serum have confirmed this instability of the normal opsonin, although even Wright himself admits that heating to 60° C. does not entirely destroy the opsonic power, though it reduces it to a minimum. A protocol from Wright and Douglas' first paper will best illustrate the degree of reduction of opsonic power resulting from the exposure of normal serum to 60-65° C. for 10 to 15 minutes.

| | |
|--|------|
| A. Unheated serum Wright—Staphylococcus suspension 1 vol.—Blood cells Wright 3 vols. | |
| (1) Phagocytic average 20 cells | 17.4 |
| (2) Phagocytic average 20 cells | 19.8 |
| B. Heated serum as above. | |
| (1) Phagocytic average 52 cells | 0.6 |
| (2) Phagocytic average 46 cells | 3.4 |

The experiments just cited refer only to the opsonic powers of normal serum. When an animal is immunized with any particular micro-organism or other cellular antigen, such as red blood cells, etc., a marked specific increase of opsonins occurs, but unlike the opsonins of normal serum these newly formed elements in the immune serum seem to possess a much greater resistance to heat.

Neufeld and Rimpau,¹⁶ who have studied these constituents of immune serum with especial thoroughness, have shown that heating to 62° to 63° C. for as long as three-quarters of an hour does not destroy them, and that such sera may be preserved for as long as several years without their complete disappearance.¹⁷

We may accept as definitely determined, therefore, that there is a qualitative difference between the serum components which initiate phagocytosis in normal serum (normal opsonins) and those which carry out the same function to a much more powerful degree in

¹⁵ Wright and Douglas. Cited in Wright, "Studien über Immun., etc.," p. 9.

¹⁶ Neufeld and Rimpau. *Deutsche med. Woch.*, No. 40, 1904; *Zeitschr. f. Hyg.*, Vol. 51, 1905.

¹⁷ Leishman. *Trans. London Path. Soc.*, Vol. 56, 1905.

immune serum. This is the more surprising since, in the case of all other antibodies (lysins, agglutinins, etc.), it has been shown that in structure and mode of action the antibodies of immune serum are in every way qualitatively similar to the corresponding ones of normal serum,^{18, 19} representing merely a specific quantitative increase of substances originally present in small amount.

This difference between the normal and immune opsonic substances has added much difficulty to the investigation of the nature of these bodies, and we may approach the problem with greater clearness by considering them separately, at first, attempting to define the relations between them after we have set down the facts ascertained in connection with each.

In their earliest investigations upon the normal opsonins Wright and Douglas²⁰ regarded them as new antibodies, separate and distinct from those already known. There is no convincing proof of this, and a number of other interpretations of the observed phenomena are possible. Indeed, the burden of proof is rather upon those who would establish the existence of a new antibody, for before this can be done it must be shown that the new function is not merely another property of the serum constituents already known. For, as Gruber has justly said, "One of the most important attributes of the natural scientist is economy of hypotheses." And in the case of the normal opsonins there are many good reasons for regarding them as possibly identical with known serum constituents. The two possibilities suggested have been (1) Are the opsonic substances identical with the alexin or complement? or (2) Do they represent the combined action of the normal sensitizer of the serum activated by the alexin?

The similarity of normal opsonin with alexin or complement has been brought out especially by Muir and Martin,²¹ by Baecher,²² and by Levaditi and Inmann.²³ The fact that both are thermolabile has been mentioned above.

In addition to this, as Muir and Martin²⁴ have shown, all antigen-antibody complexes which absorb alexin out of serum at the same time remove the normal opsonin. Thus sensitized red corpuscles, sensitized bacteria, and specific precipitates added to normal serum take out its opsonic substances. From this fact they also concluded that the normal opsonins like alexin were non-specific. For just as

¹⁸ Dean. *Proc. Royal Soc.*, 76, 1905.

¹⁹ Neufeld and Hüne. *Arb. a. d. kais. Gesundh. Amt.*, Vol. 25, 1907.

²⁰ Wright and Douglas. *Loc. cit.*

²¹ Muir and Martin. *Br. Med. Jour.*, Vol. 2, 1906; *Proc. Royal Soc. B.*, Vol. 79, 1907.

²² Baecher. *Zeitschr. f. Hyg.*, Vol. 56, 1907.

²³ Levaditi and Inmann. *C. R. de la Soc. Biol.*, 1907, pp. 683, 725, 817, 869.

²⁴ Muir and Martin. *Loc. cit.*

the alexin of a serum may serve to activate a considerable variety of sensitized antigens, so the opsonic action of a normal serum may functionate upon a large variety of bacteria. Muir and Martin were probably wrong in this and, as we shall see below, normal opsonins, like normal sensitizers, may be regarded as specific.

Similar to the observations of Muir and Martin are those of Neufeld and Hüne,²⁵ which showed that yeast cells will absorb both alexin and opsonin out of serum.

A further similarity between the two serum constituents is the fact that both are absent from the normal fluid of the anterior chamber of the eye, but they together appear in it after injury (puncture for the first removal of fluid). A like parallelism between the absence and presence of both has been shown for edema fluids. Furthermore, phosphorus poisoning which reduces alexin likewise reduces opsonin.

Although this parallelism is very striking, it does not on this account mean that necessarily the two are identical. It may signify merely that the alexin is a necessary participant in normal opsonic action, essential in that it activates a thermostable opsonic constituent just as it activates hemolytic or bactericidal sensitizer.

This opinion has been expressed by Levaditi, Neufeld,²⁶ Dean,²⁷ and others, and indeed it is a conception which seems most logical. For the procedures which remove both alexin and opsonin, as stated above, do not, as a matter of fact, remove *all* the opsonic action. (Although Neufeld maintains this.²⁸) Studies of Hektoen and others have definitely proved that, though reduced to almost nil, nevertheless heated serum shows definite though slight opsonic action as compared with indifferent menstrua such as salt solution. A similar slight remnant of opsonic action after absorption of normal serum with sensitized cells, bacteria, and precipitates is evident in the protocols of Muir and Martin. The significance of this point becomes immediately clear when we consider the properties of the bacteriotropins or immune opsonins, which are heat stable and capable of initiating opsonic action in the entire absence of alexin or complement. It is possible, therefore, that there may be present in normal serum a slight amount of specific thermostable opsonin, which, though capable of acting feebly by itself, is nevertheless powerfully activated by alexin—just as bactericidal or hemolytic antibody is similarly activated.

One of the most thorough studies upon this question is that of

²⁵ Neufeld and Hüne. *Arb. a. d. kais. Gesundh. Amt.*, Vol. 25, 1907.

²⁶ Neufeld. "Kolle u. Wassermann's Handbuch," *Ergänzungsband* 2, p. 313.

²⁷ Dean. *Brit. Med. Jour.*, 2, 1907, p. 1409.

²⁸ In fact he states that heated normal serum may be used as a control in opsonic experiments instead of salt solution.

Cowie and Chapin.²⁹ Dean³⁰ had previously shown that, although heated immune serum was capable of exerting opsonic action by itself, this action could nevertheless be enhanced by the addition of a little diluted fresh normal serum. The particular significance of Dean's work will be discussed later. Cowie and Chapin, however, carried on similar experiments with normal serum in which they attempted to reactivate heated normal serum by the addition of small amounts of diluted fresh serum, by itself but slightly opsonic. One of their experiments may serve to illustrate this point, as follows:

Experiment 10. June 13, 1907

| | Phagocytic count ³¹ |
|--|-----------------------------------|
| 1. Unheated serum..... | 15.44 |
| 2. Salt solution..... | 0.18 |
| 3. Heated serum, 57° C..... | 1.08 |
| 4. Diluted serum (1:15)..... | 1.56 |
| 5. Heated serum 57° C. + diluted serum (1:15)..... | 12.40 |
| 6. Unheated serum + unheated serum..... | 16.08 |

This experiment and others like it seem to demonstrate clearly that the opsonic action of normal serum, though dependent largely upon alexin, is nevertheless also dependent upon a heat-stable body, comparable to the sensitizer or amboceptor, in that it is reactivable to almost the full power of the original condition (before heating) by slight amounts of alexin—in themselves almost inactive.³²

These findings were later confirmed by Eggers,³⁴ and it is plain from this work that the apparent opsonic inactivation of normal serum by heat depends upon the destruction of the heat-sensitive constituent only—the heat-stable substance—surely involved in the process, remaining intact, and reactivable.

Closely associated with this phase of the problem is that of the specificity of the normal opsonins. For if, as at first supposed, the normal opsonins are, like complement or alexin, non-specific, the above amboceptor-complement structure of this mechanism would be rendered unlikely. Earlier work upon this question was contradictory. Bulloch and Western,³⁵ working with staphylococci and

²⁹ Cowie and Chapin. *Jour. Med. Res.*, Vol. 17, 1907, pp. 57, 95 and 213.

³⁰ Dean. *Loc. cit.*

³¹ Phagocytic count = average number of bacteria in each leukocyte.

³² In earlier experiments Hektoen and Ruediger³³ did not succeed in reactivating heated sera and concluded that normal opsonins had the hypothetical structure of toxins in that they possessed a haptophore and an opsonophore group. From this point of view Hektoen has subsequently receded largely because of work done under his own direction.

³³ Hektoen and Ruediger. *Jour. Inf. Dis.*, 1905.

³⁴ Eggers. *Jour. of Inf. Dis.*, Vol. 5, 1908.

³⁵ Bulloch and Western. *Proc. Roy. Soc. B.*, 77, 1906.

tubercle bacilli, found that each of these organisms absorbed out separately specific opsonins from normal serum, leaving those for other bacteria but slightly reduced. Slight reduction of the opsonic action for other micro-organisms might easily be explained by a partial removal of complement which is bound to take place in such experiments. Simon, Lamar and Bispham,³⁶ and some others failed to find any such specificity. Russell,³⁷ Axamit and Tsuda,³⁸ and a number of others obtained similar negative results—in that a number of different bacteria seemed to absorb opsonins out of normal serum indiscriminately and without specificity. On the other hand, more recent careful work by Rosenow,³⁹ by Macdonald,⁴⁰ and by Hektoen⁴¹ has upheld the original contention of Bulloch and Western. The work of Rosenow, in which pneumococci were shown to absorb out their specific opsonins from normal human serum, taking out in part only those for streptococci, staphylococci, and tubercle bacilli, is especially convincing, and the experiment of Hektoen with normal hemopsonins (opsonins which cause the phagocytosis of red blood cells) bear him out.

It seems fair to conclude, therefore, that normal opsonins depend upon the coöperation of a heat-stable and a heat-sensitive body. The heat-stable body, analogous to normal sensitizer or amboceptor, is specific and reactivable by the heat-sensitive body which appears to be identical with alexin. This statement merely asserts the facts of the dual mechanism of the process without assuming necessarily the identity of the heat-stable body with sensitizer or that of the heat-sensitive one with alexin, though this seems extremely probable.

This question we will discuss again more particularly in connection with the bacteriotropins or immune opsonins.

Further proof for such a complex constitution of the normal opsonins has been adduced by means of absorption experiments at 0° C.—by Cowie and Chapin. In our discussion of the lytic antibodies we have seen that sensitizer or amboceptor may be absorbed from serum by its specific antigen at 0° C.—but that the attachment of alexin takes place only when the temperature is raised above this. Practically no alexin is bound at the low temperature. Cowie and Chapin, applying this method of investigation, showed:

1. That normal human serum may have its opsonic power for staphylococci removed by absorption with staphylococci at 0° C.
2. Serum so treated retains the power of reactivating the opsonin of heated normal serum.

³⁶ Simon, Lamar, and Bispham. *Jour. Exp. Med.*, Vol. 8, 1906.

³⁷ Russell. *Johns Hopk. Bull.*, Vol. 18, 1907.

³⁸ Axamit and Tsuda. *Wien. klin. Woch.*, Vol. 20, No. 35, 1907.

³⁹ Rosenow. *Jour. Inf. Dis.*, Vol. 4, 1907.

⁴⁰ Macdonald. Quoted from Hektoen, *loc. cit.*; *Aberdeen Univ. Studies*, Vol. 21, 1906, p. 323.

⁴¹ Hektoen. *Journ. Inf. Dis.*, Vol. 5, 1908.

3. Staphylococci so treated are more easily subject to phagocytosis in the presence of dilute normal serum, or normal serum which has been inactivated by contact with staphylococci in the cold, than are the same bacteria untreated.

Kurt Meyer⁴² has carried out similar experiments with paratyphoid bacilli and normal serum, and, though his work is less extensive, he reaches the same conclusion as Cowie and Chapin.

We may accept, therefore, as fairly well established that the opsonic power of normal serum depends upon a complex mechanism consisting of (a) a thermostable substance comparable to amboceptor or sensitizer, probably specific, but present in very small amount, and (b) a thermolabile substance probably identical with alexin or complement which powerfully, but non-specifically, enhances the slight opsonic power of the thermostable substance.

In considering this conception, together with the subsequent discussion of bacteriotropins or immune opsonins, it will be well to remember that in normal inactivated sera the thermostable opsonic constituent differs in its action from the bodies we speak of as amboceptors or sensitizers in that it may functionate for phagocytosis by itself—entirely without alexin—while neither bactericidal nor hemolytic effects can be brought about by sensitizer alone. Does this definitely exclude the identity of this thermostable opsonic substance and sensitizer? It is indeed an argument against identification, but in opsonic action, we must remember, there is merely a sensitization to the action of the phagocyte. This phagocyte may in itself be capable of furnishing a small amount of substance comparable in action to alexin—in fact, we have seen that the origin of alexin from leukocytes is still suspected by a number of workers. At any rate the phagocyte is a living cell which may well be capable of supplying in itself to some degree the necessary activation, and therefore the difference cited above is not necessarily a proof that the normal thermostable opsonic constituent is different from normal sensitizer or amboceptor.

The difference between the opsonic action of normal serum and that of immune serum, then, is the fact that heating to from 56° to 60° C. almost completely destroys the former, whereas it has but slight if any diminishing effect upon the latter. The immune opsonins, or, as Neufeld and Rimpau have called them, bacteriotropins, therefore are thermostable. This was determined as early as 1902 by Sawtschenko,⁴³ and was subsequently studied with great accuracy

⁴² Kurt Meyer. *Berl. klin. Woch.*, 1908, p. 951.

⁴³ Sawtschenko. *Ann. de l'Inst. Past.*, Vol. 16, 1902, quoted from Levaditi.

by Neufeld and Rimpau,⁴⁴ Neufeld and Töpfer,⁴⁵ Dean,⁴⁶ Hektoen,⁴⁷ and others. It was shown that when an animal is immunized with any given bacterium or other cellular antigen (blood corpuscles, etc.) opsonic substances specific for the particular antigen appear in considerable quantities, and these are but slightly, if at all, diminished when the serum is heated; Neufeld and Hüne⁴⁸ found that heating for as long as three-quarters of an hour to 63° C. did not noticeably reduce the activity of the bacteriotropins of immune serum, and that, again, unlike the normal opsonins, prolonged preservation, under sterile conditions, changes them but slowly.

These facts alone indicate a close similarity between the bacteriotropins and the other well-known thermostable constituents of immune sera, and the question here again immediately arises whether we are to regard them as identical with any of the other specific antibodies or as distinct substances independent of these.

It was suggested early in these investigations by Muir and Martin that bacteriotropins might be identified with agglutinins, inasmuch as they possessed resistance to heat, were active without apparent dependence upon alexin, and could not, at least in the earlier studies, be reactivated by the addition of fresh normal serum when once inactivated. The supposition was that for this reason the bacteriotropin might have a structure like the hypothetical "haptines of the second order" which Ehrlich attributes to the agglutinins. This supposition has found no experimental support in that agglutination and bacteriotropic effects did not run parallel. We ourselves are not ready to admit that such lack of parallelism is proof against their identity. However, since it is very probable that both agglutination and precipitation are merely phenomena of colloidal flocculation effects which follow certain quantitatively adjusted combinations of antigen and specific antibody, and that it is not at all necessary to assume separate agglutinating or precipitating serum constituents, this problem becomes merely another version of the question of the identity of bacteriotropins and sensitizer or amboceptor.

Apart from thermostability, further similarity lies in the fact that bacteriotropins are strictly specific and may be specifically absorbed out of immune sera by their respective bacteria.

Like amboceptor or sensitizer they are specifically increased to a powerful degree by the treatment of animals with any given microorganism and may be incited not only by the injection of bacteria

⁴⁴ Neufeld and Rimpau. *Deutsche med. Woch.*, No. 40, 1904; *Zeitschr. f. Hyg.*, 51, 1905.

⁴⁵ Neufeld and Töpfer. *Centralbl. f. Bakt.*, 1, 38, 1905.

⁴⁶ Dean. *Proc. Roy. Soc. B.*, 76, 1905.

⁴⁷ Hektoen. *Jour. Inf. Dis.*, 3, 1906, and *loc. cit.*

⁴⁸ Neufeld and Hüne. *Arb. a. d. kais. Gesundh. Amt.*, Vol. 25, 1907.

but by that of blood cells as well. In spite of these points of likeness, however, Neufeld ⁴⁹ and his associates maintain rigidly that the two substances are not the same and that the bacteriotropins are distinct and independent antibodies.

Among the reasons advanced in support of this opinion are the facts that certain immune sera, both antibacterial and hemolytic, may contain bacteriotropins without containing lysins and vice versa. That this is undoubtedly true has been shown not only by Neufeld and his associates but by Hektoen ⁵⁰ and others, and it is likewise a fact that in sera in which both functions are demonstrable they frequently do not run quantitatively parallel. These are unquestionably strong arguments, but their force is somewhat weakened, as Levaditi has pointed out, by the fact that there are many varieties of bacterial immune sera which undoubtedly sensitize the specific bacteria (as can be shown by alexin fixation), but which do not lead to bacteriolysis. Wassermann ⁵¹ also attaches little value to the lack of parallelism between the lytic and opsonic functions, expressing the belief that the solubility of the particular antigen may determine whether sensitization leads to phagocytosis or to lysis. With bacteria like the cholera spirillum rapid lysis takes place, but when, as in pneumococci or streptococci, there is great resistance to lysis, sensitization may lead to delayed lysis anticipated by leukocytic accumulation, phagocytosis, and intracellular digestion.

It by no means follows from mere lack of parallelism, therefore, that the two serum functions are dependent upon separate antibodies, although the argument is sufficiently strong to impose conservatism in identifying them.

Another important argument advanced against the identification of bacteriotropins with the bactericidal sensitizers or amboceptors is the fact that the former lead to phagocytosis without the participation of alexin, whereas the latter become active for lysis only when alexin is present.

This point also has constituted Neufeld's strongest support for maintaining that the bacteriotropins or immune opsonins are entirely distinct from the normal opsonins. It is true, indeed, that immune serum, unlike normal serum, may opsonize powerfully even after heating to temperatures which destroy alexin.

If we regard the heat-stable lytic antibody as an amboceptor in the strict sense of Ehrlich, as a specific "Zwischenkörper" with a complementophile group, this argument would have considerable weight. Even in this case, however, strong sensitization of the bacteria may make them amenable to the living cells—the phagocytes—

⁴⁹ Neufeld and Töpfer. *Centralbl. f. Bakt.*, 1, 38, 1905.

⁵⁰ Hektoen. *Jour. of Inf. Dis.*, 6, 1909.

⁵¹ Wassermann. *Deutsche med. Woch.*, Vol. 33, No. 47, 1907.

which in itself may furnish a slight amount of alexin or alexin-like substances.

We may regard the action of the immune serum upon the antigen as rather a sensitization in the sense of Bordet, and it does not seem logical to assume that the heat-stable bodies, similar in other respects, are different merely because they can sensitize bacteria both to the action of an alexin and to that of a living cell, which in itself surely contains a number of different enzymes, comparable functionally to alexin, though possibly not identical with it.

Indeed, the experiments of Dean have given much positive evidence in favor of regarding the immune opsonins or bacteriotropins as true amboceptors or sensitizers. Dean⁵² found that, although heated immune serum may unquestionably opsonize by itself, its action may be still further enhanced by the addition of a little diluted normal serum (compare these results with those of Cowie and Chapin on normal opsonins). Hektoen's⁵³ experiments with hemopsonic immune sera are analogous. We cite one of these as illustrating the point in question:

TABLE I

Phagocytosis of Goat Corpuscles under the Influence of Goat-blood-immune Rabbit Serum, and Normal Guinea Pig Complement (Table from Hektoen, loc. cit.)

| Immune serum | | Normal guinea pig serum | Phagocytosis |
|--------------|---|-------------------------|--------------|
| .001 | | — | 4. |
| .001 | + | .01 | 20. |
| | | .01 | 0 |

Here, therefore, the diluted immune serum, but slightly cytotropic in itself, was powerfully activated by a diluted unheated normal serum, which in itself was entirely inactive.

Indeed, an experiment by Neufeld himself, with Bickel,⁵⁴ points in the same direction. They found that, when a heated specific hemolytic serum was added to the homologous cells in such small quantities that it no longer exerted cytotropic (opsonic) action, the addition of a small amount of alexin, too small to lead to hemolysis of the cells (and not by itself cytotropic or hemopsonic), caused active phagocytosis. Analogous experiments upon bacterial antisera were carried out by Levaditi and Inmann. It thus appears that, even in the case of the immune opsonins or bacteriotropins, we may think of a participation of two substances—a sensitizer-like one and one comparable to alexin or complement. We may, at least, infer that the full opsonic action both of normal and immune sera is dependent

⁵² Dean. *Proc. Royal Soc. B.*, 79, 1907.

⁵³ Hektoen. *Jour. Inf. Dis.*, Vol. 6, 1909, p. 67.

⁵⁴ Neufeld and Bickel. *Arb. a. d. kais. Gesundh. Amt.*, Vol. 27, 1907.

upon the coöperation of two such bodies. It is likely, therefore, that the mechanism of normal and of immune opsonic action may, after all, differ only in quantitative relations between the two.

For assuming this to be an antibody-alexin mechanism like hemolysis, we may recall the work of Morgenroth and Sachs on the relations between amboceptor and complement in hemolysis. There we saw that a large amount of amboceptor would cause hemolysis in the presence of a small amount of complement and vice versa. Therefore, here, too, in normal serum the small quantity of amboceptor or specific thermostable opsonin (bacteriotropin) may act very powerfully in the presence of the alexin. When the latter is destroyed, however, the minute quantity of specific thermostable opsonin is hardly enough to do more than initiate slight phagocytosis of comparatively non-resistant bacteria, whereas the large amount of specific sensitizer left in immune sera after inactivation may still lead to strong bacteriotropic action. In outlining this explanation we have consistently drawn upon the analogy between thermostable opsonin and amboceptor or sensitizer. Whether or not these two substances are identical is by no means positively determined and must be considered an open question for the present. However, from the above, it seems to us that much testifies in favor of such an identification.⁵⁵

The preceding discussions have ignored the possibility that apart from opsonic or bacteriotropic action on the bacteria there may be a difference in phagocytic energy which depends upon inherent properties of the leukocyte itself.

Indeed, the technique by which the researches of Wright and his followers were carried out does not in any way take into account the source of the leukocytes as a possible variable factor. There is, however, a considerable amount of evidence which points to differences in phagocytic powers residing in the leukocytes themselves independent of the serum. Park and Biggs⁵⁶ have demonstrated such differences for the leukocytes of normal persons in the phagocytosis of staphylococci, and more extensive researches have been made with similar results, in the case both of staphylococci and tubercle bacilli by Glynn and Cox.⁵⁷

The last-named authors, moreover, recognized the necessity, in making such investigations, of experimenting with leukocyte emulsions containing approximately the same number of cells, for, as Fleming⁵⁸ had shown, if unequal leukocytic emulsions are used,

⁵⁵ Pfeiffer (quoted from P. Th. Müller) regards opsonic action as due to a combined action of amboceptor and complement and speaks of it as an "Andauung" of the bacteria for the leukocyte—which we may translate best as a partial predigestion.

⁵⁶ Park and Biggs. *Jour. Med. Res.*, Vol. 17, 1907.

⁵⁷ Glynn and Cox. *Jour. Path. and Bact.*, 14, 1910.

⁵⁸ Fleming. *Practitioner*, London, Vol. 80, 1908.

less phagocytosis per cell occurs in the emulsion containing the greater number of leukocytes. This phase of the subject has been taken up most thoroughly by Hektoen⁵⁹ and his associates, and Rosenow⁶⁰ has made careful comparative studies on pneumococcus phagocytosis, in which he standardized the leukocytic suspensions by actual cell counts. His work as well as that of Tunnicliff,⁶¹ of the same school, has shown definitely that the inherent phagocytic power of leukocytes may vary not only in health and disease, but differences may exist between the cells of apparently normal people. Tunnicliff showed, for instance, that at birth the leukocytes are less active than in adult life.

For accurate experimental work, therefore, as well as in theoretical reasoning upon problems of phagocytosis, it is necessary to bear in mind the possible inherent variations in the leukocytes themselves.

Of the three factors concerned in the process of phagocytosis, then, we have considered two, the serum and the leukocytes. The former we have seen exerts a powerful determinative influence on the process, the latter a less marked influence, though still definite and measurable. We have still to discuss the bacteria themselves as variable factors in determining the degree to which phagocytosis may take place.

This problem was first investigated by Denys and Marchand in connection with their work upon streptococcus immunity, and was further studied in detail by Marchand. Marchand⁶² showed that leukocytes would readily take up non-virulent streptococci in the presence of normal serum, but that under similar conditions virulent streptococci were not phagocytosed at all or to a very slight degree only. He determined further that this resistance to phagocytosis remained unchanged after the virulent organisms had been killed by heat, and washed clean of culture fluid. It seemed, therefore, that the resistance depended upon a condition of the bacterial body and not upon substances secreted and given off to the environment. These experiments, as well as similar work by Mennes,⁶³ Gruber and Futaki,⁶⁴ and others makes it clear that differences in virulence between different species of bacteria, as well as between different strains of the same micro-organism, depend, at least in part, upon the resistance which the bacterial bodies oppose to ingestion by the leukocytes. We must distinguish clearly here between these apparently purely "antiopsonic" bacterial properties and those supposedly "antichemotactic" substances which are conceived as a cause for

⁵⁹ Hektoen. *Jour. of A. M. A.*, Vol. 57, No. 20, 1911.

⁶⁰ Rosenow. *Jour. of Inf. Dis.*, 7, 1910.

⁶¹ Tunnicliff. *Jour. Inf. Dis.*, 8, 1911.

⁶² Marchand. *Arch. de Méd. Exp.*, No. 2, 1898.

⁶³ Mennes. *Zeitschr. f. Hyg.*, Vol. 25, 1897.

⁶⁴ Gruber and Futaki. *Münch. med. Woch.*, 1906.

virulence by Deutsch and Feistmantel⁶⁵ and by Bail⁶⁶ in his so-called "aggressins." The latter are supposed to be secreted bacterial substances by means of which the leukocytes are held at bay. The properties we are, at present, considering are probably in no way antichemotactic, but oppose purely the actual ingestion by the leukocyte, nor do they seem to depend upon the secretion of substances which injure the leukocytes. For, in the first place, a profuse accumulation of leukocytes may follow the injection of virulent micro-organisms, and Denys (quoting from Gruber) has seen active phagocytosis of virulent pneumococci, but none of virulent streptococci when antipneumococcus serum was injected with the mixture.

Rosenow⁶⁷ has carried out a thorough investigation dealing with these relations in pneumococcus infection. Seventy-five strains of this organism were all found non-phagocytatable when first isolated and the resistant condition was associated with virulence for rabbits and guinea pigs. It was found, moreover, that the resistance to phagocytosis was dependent upon the inability to absorb opsonin. For, while phagocytatable non-virulent pneumococci absorbed specific opsonin from serum, the virulent ones failed to do this in proportion to the degree of their virulence. Furthermore, extraction of the bodies of the virulent organisms in NaCl solution yielded a substance which inhibited the action of pneumococcus opsonin—a true anti-opsonin—which he speaks of as "virulin." This discovery, if confirmed, would supply us with a very simple explanation for some phases of the problem of virulence. It is, indeed, likely that the antiopsonic property is closely bound up with chemical and structural changes which take place in the bacterial cell as it adapts itself to the parasitic conditions. This is plain from the fact that pneumococci and some other bacteria will rapidly lose their virulence when cultivated on artificial media devoid of animal serum, will retain it longer if grown on some serum media, and will rapidly regain it if passed through animals. The formation of a capsule is unquestionably a morphological evidence of such a change. Habitually capsulated bacteria, like the Friedlander bacillus, and *Streptococcus mucosus*, are of fairly constant virulence, while in other micro-organisms like the pneumococci, anthrax bacillus, plague bacillus, and certain other streptococci, the formation of a capsule goes hand in hand with an increase of virulence. By the aid of this morphological earmark of virulence, moreover, Gruber and Futaki have obtained further

⁶⁵ Deutsch and Feistmantel. Quoted from Sauerbeck. *Lubarsch und Ostertag*, Vol. 2, 1906.

⁶⁶ Bail. *Arch. f. Hyg.*, Vol. 52, 1905.

⁶⁷ Rosenow. *Jour. Inf. Dis.*, Vol. 4, 1907.

proof that the resistance to phagocytosis in these cases is due to the nature of the bacterial cell body rather than to any secreted anti-opsonic substances. For, after the injection of anthrax bacilli into guinea pigs, they saw that leukocytes would take up unencapsulated bacilli, apparently picking them out of the midst of surrounding encapsulated organisms which they were unable to ingest.

CHAPTER XIV

THE OPSONIC INDEX AND VACCINE THERAPY

WRIGHT'S¹ investigations upon phagocytosis were, indirectly, the outcome of his earlier work upon antityphoid vaccination. His purpose in these studies had been a purely practical one, and he had attempted to obtain a guide for the dosage and the interval between injections by measuring the bactericidal and agglutinating powers of the blood serum. In the case of typhoid immunization this was indeed a practicable method of control, since the bactericidal power of the blood serum rose directly as the immunization of the patient was attained. In the cases of many other bacteria, however, this method of study was not practicable, and Wright, as others before him, did not find a regularly increased specific bactericidal power in the blood sera of immunized animals or of patients convalescing from infections with such bacteria as the staphylococcus, streptococcus, *Micrococcus melitensis*, the *Bacillus pestis*, and a number of others. In fact, together with Windsor,² he showed that normal human blood has practically no bactericidal power for pyogenic staphylococci and that antistaphylococcus inoculations or recovery from an infection do not result in the production of such properties in the serum. These determinations are practically identical with Nuttall's³ earlier studies on the same bacteria and, indeed, correspond with the data obtained by Metchnikoff and his followers in their work on anthrax infection. For, in discussing these investigations, we saw that very often the serum of a comparatively resistant animal is less potently bactericidal than that of a more susceptible one. We need only recall the difference between rabbits and dogs in this respect. The serum of the former is more strongly bactericidal than that of the latter, and yet rabbits are the far more susceptible animals. These relations have been studied with great care, also, by Petterson.⁴ It was logical in such cases to look for the cause of resistance in the activity of the phagocytes, and this, we have seen, Metchnikoff did successfully in a large series of cases, both as regards natural and acquired immunity.

¹ Wright. *Lancet*, 1902; *Practitioner*, Vol. 72, 1904.

² Wright and Windsor. *Jour. of Hyg.*, Vol. 2, 1902; and Wright, *Lancet*, 1900 and 1901.

³ Nuttall. *Zeitschr. f. Hyg.*, Vol. 4, 1888.

⁴ Petterson. *Centralbl. f. Bakt.*, Vol. 39.

Yet the controversy between the strictly humoral and the cellular schools was by no means regarded as closed, especially since, in such cases as typhoid infection, the parallelism between increased resistance and extracellular bactericidal power was so plainly evident, while in this disease particularly (for technical reasons which will become clear as we proceed) no such parallelism with phagocytosis could at first be shown. It was because of such apparent confusion that Leishmann⁵ undertook to study again the relation of phagocytosis to active immunity, chiefly upon staphylococcus cases that were being "vaccinated" therapeutically by Wright himself.

In order to obtain a numerical measure of the degree of phagocytosis, he developed a simple technique which, though crude, served to give him the information he sought. It consisted in taking small quantities of the blood of patients and mixing these in capillary pipettes with equal volumes of bacteria suspended in salt solution.

The mixtures were then placed on slides, covered with a coverslip, and incubated at 37° C. for varying periods. At the end of incubation the preparations were smeared upon slides and stained by Leishmann's modification of the Romanowski method, the number of bacteria in a large series of leukocytes counted and an average taken.

This method had many serious flaws, chief among them being the liability to coagulation of the preparations and the fact that, in each test, the fluid constituents as well as phagocytes, both of them variable factors, came from the same individual. While, therefore, it was possible to estimate an increase or decrease of general phagocytic power, it was impossible to analyze this in reference to its dependence either upon the condition of the cells, on the one hand, or that of the plasma or serum, on the other. Moreover the relation of the number of leukocytes to that of bacteria in individual tests necessarily differed, and this, we have seen, adds a variable factor which renders it impossible to compare any two experiments with accuracy.

In spite of these difficulties, however, Leishmann succeeded in establishing, in a number of cases of staphylococcus infection, that an increased resistance was accompanied by an increased energy of phagocytosis.

Leishmann, however, went no further than this, and interpreted his results on the basis of the "stimulin" theory of Metchnikoff.

The subsequent studies of Wright, which began at the point at which Leishmann stopped, have been described in the preceding chapter and had, as their main result, we have seen, the discovery of the opsonins and the final confirmation of Denys' conception of the true mechanism of coöperation between serum and leukocytes in phagocytosis. In order to carry out these studies the technique of Leish-

⁵ Leishmann. *Br. Med. Jour.*, 1, 1902; *Transact. Lond. Path. Soc.*, Vol. 56, 1905.

mann was quite inadequate, and Wright's first task was to modify it in such a way that reasonably accurate comparative estimates of phagocytosis could be made.

It is necessary to outline Wright's method briefly in this place in order that we may consider possible sources of error and obtain a clear understanding of the conclusions he based on his observations.



WRIGHT CAPSULE FOR TAKING BLOOD TO OBTAIN SERUM FOR OPSONIC TESTS.

Wright recognized that the determination of the degree of phagocytosis, induced by the opsonin of any given serum in a single test, is by itself of no value, since the actual number of bacteria taken up by each leukocyte, apart from the opsonic contents of the serum, depends also upon such purely technical factors as the concentration of the bacterial emulsion, the relative number of leukocytes, and the length of time of incubation. Two individual tests, therefore, carried out with the serum of the same patient at the same or at different times, with different bacterial emulsions or leukocytes in each, would give variable results, even though the opsonin contents themselves were entirely alike.

In order, therefore, to obtain a relative estimate of the opsonic contents of any serum it is necessary to compare the phagocytic activity induced by this serum with the similar power of another supposedly normal serum, both tests being carried out, under exactly similar conditions, with the same bacterial emulsion and the same leukocytes. The average number of bacteria found in each leukocyte in each one of the preparations is then the "phagocytic index." The relation of the phagocytic index of the unknown serum to that of the supposedly normal serum constitutes what Wright has called the "opsonic index."

Instead of using the whole blood of the patient Wright takes a small amount of blood in glass capsules, allows it to clot, and uses the expressed serum in his test. For comparison with this he employs a "pool" of a number of specimens of serum from supposedly

Wright recognized that the determination of the degree of phagocytosis, induced by the opsonin of any given serum in a single test, is by itself of no value, since the actual number of bacteria taken up by each leukocyte, apart from the opsonic



METHOD OF PRODUCING AN EVEN EMULSION OF BACTERIA FOR OPSONIN DETERMINATION.

normal individuals. By the use of such a serum mixture any slight possible variations from the normal in any one of the sera are likely to be equalized, and a closer approach to a normal standard is attained.

The leukocytes used in both tests are the same and taken, as a rule, from the blood of the worker or from some other supposedly healthy person. They are obtained by taking 15 or 20 drops of blood from the finger or ear into 5 to 10 c. c. of sodium citrate solution, in which the blood does not clot. Brief centrifugalization throws down the blood cells, with a thin, buffy coat of leukocytes on top, and these are gently taken off with a pipette. This constitutes the leukocytic cream of Wright's experiments, and furnishes a uniform leu-



METHOD OF TAKING UP EQUAL VOLUMES OF LEUKOCYTES, BLOOD SERUM AND BACTERIAL EMULSION IN WRIGHT'S TECHNIQUE FOR OPSONIC-INDEX DETERMINATION.

kocyte factor for the two tests which are to be compared. The bacteria are obtained by emulsifying carefully in salt solution. It is very important to obtain an emulsion free from clumps and neither too thick nor too thin, a result which can be secured only by experience.

Equal quantities of serum (unknown and normal "pool" respectively) are mixed with equal quantities of the bacterial emulsion and the leukocytes in capillary pipettes, and the mixtures are incubated for fifteen to thirty minutes under exactly similar conditions. At the end of this time smears are made upon slides, the preparations stained, and the numbers of bacteria in a hundred or more leukocytes counted in each of the two experiments. The average is taken, and from the *phagocytic indices* thus obtained the *opsonic index* is calculated. For instance, if

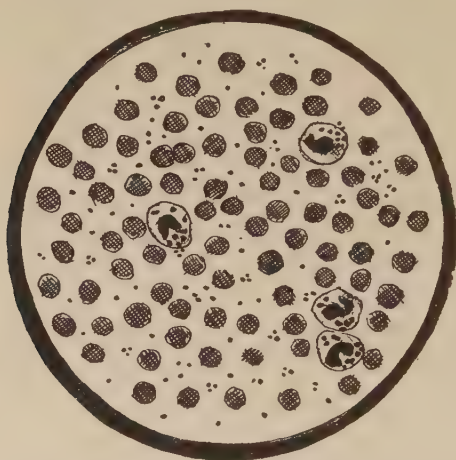
$$\begin{aligned}\text{Phagocytic index (normal pool)} &= 8 \\ \text{Phagocytic index (patient's serum)} &= 6\end{aligned}$$

then the *opsonic index* (patient's serum) = 0.75. Or, if the phagocytic index of the normal pool had been 10. and that of the patient's serum 15., then the *opsonic index* of the patient's serum, higher than normal, would be 1.5.

For the insurance of accuracy in carrying out this method Wright calls especial attention to the caliber of the capillary pipettes that are used, the concentration of the sodium citrate solution, which should be 1.5 per cent., and the freshness of the leukocytes. But it is still necessary to remember that with the greatest care in tech-

nique uncontrollable sources of error influence this method. Most important among them are the differences necessarily existing between different normal sera used for comparison and differences in the agglutinative powers of the sera used in the two specimens. For it is plain that different degrees of agglutination may bring about great variations in the number of bacteria with which the individual leukocyte comes into contact.

Wright's method has also been particularly unsatisfactory in taking the opsonic index against such bacteria as the typhoid bacillus



LEUKOCYTES CONTAINING BACTERIA. DRAWING OF FIELD AS SEEN IN WRIGHT'S METHOD OF OPSONIC-INDEX ESTIMATION.

and the cholera spirillum, organisms which are very rapidly digested after being taken up by the leukocytes. In consequence, even after as short an incubation time as five or ten minutes, the ingested bacteria are partly disintegrated, are stained indistinctly, and cannot be counted with accuracy. In order to avoid this source of error Klien⁶ has devised a modification which depends upon gradual dilution of the serum in a series of phagocytic tests with the same leukocytic and bacterial emulsions. In this way he determines the degree of dilution of the serum to be

tested at which phagocytosis no longer exceeds that taking place in salt solution alone. The degree of dilution at which this result was obtained has been called by Simon the "coefficient of extinction." A comparison of sera with regard to this value, it is clear, furnished an estimate of their quantitative opsonic properties quite as instructive as the direct estimations by the Wright method, and in our opinion, at least, more reliable. Though also subject to some of the objections advanced against the Wright method, it has the definite advantages mentioned above, and is not so closely dependent upon irregularities in counting, agglutinin influences, and differences in relative proportions of bacteria and leukocytes employed. Jobling⁷ has used this method with success for the standardization of antimeningitis serum.

A further modification suggested by Simon, Lamar, and Bis-

⁶ Klien. *Johns Hop. Hosp. Bull.*, Vol. 18, 1907.

⁷ Jobling. "Studies from the Rockefeller Inst.," Vol. 10, 1910, p. 614.

pham⁸ depends upon a combination of the dilution method and a modification in the method of counting. They make comparative tests of the same serum, diluted from 1 to ten to 1 to one hundred in salt solution, and estimate the opsonic power, not by determining the average number of bacteria to the leukocyte, but by taking a percentage of the total number of leukocytes which take part in the phagocytosis, that is, contain any leukocytes at all. The bacterial emulsion for this method should be so thin that, in normal serum, only about 50 per cent. of the leukocytes will contain bacteria.

That Wright's method, or any of the others, gives absolutely accurate results will hardly be claimed by any one who has worked upon opsonic-index estimations. There are certain uncontrollable variable factors, some of which have been pointed out above; and, apart from these, the delicacy of the technique is such that reliable results can ordinarily be obtained only by trained workers after considerable practice and experience. Even in such hands the percentage of personal error is more likely to be above than below 10 per cent. For ordinary clinical purposes, therefore, in the control of cases the estimation of the opsonic index is not often practicable.

On the other hand, there can be little doubt about the fact that careful comparative estimation, by Wright's method and by some of the modifications, carried out by workers with experimental training and consequent attention to extensive controls, have yielded results of sufficient accuracy to permit the recognition of definite facts concerning opsonins. It is beyond question, therefore, that the conclusion regarding the relation of opsonic fluctuations to clinical conditions and the general significance of opsonins emanating from laboratories like those of Wright, Neufeld, Hektoen, and some others may be accepted as fact—especially since in most essentials such workers have agreed. In consequence we are now in possession of knowledge regarding the opsonic constituents of the blood in health and disease, and in the course of active immunization with bacterial vaccines, which is of the greatest practical importance. We may summarize the results of such investigations by saying that in many of the infections of man the resistance of the patient is roughly proportionate to the opsonic index—and that properly spaced inoculation with suitable quantities of dead bacteria (vaccines) will raise the opsonic index and lead to recovery in many of the localized subacute and chronic conditions.

As to the usefulness of the treatment in various infections and the limitations within which we may hope for results opinions differ, and these will be discussed more fully below. Before we proceed to this, however, it will be useful to consider the studies upon which

⁸ Simon and Lamar. *Johns Hop. Hosp. Bull.*, Vol. 17, 1906; Simon, Lamar, and Bispham, *Jour. Exp. Med.*, Vol. 8, 1906; Simon, *Jour. A. M. A.*, Vol. 48, 1907, p. 139.

the parallelism between opsonic index and clinical condition was founded.

Wright's own earlier studies were made chiefly upon staphylococcus infections and tuberculosis. Since then the method has been applied to almost all known infections with varyingly successful results.

One of the first steps in determining such a parallelism between the resistance of a patient and the opsonic index consisted, of course, in comparing the index of the sera of normal individuals with that of patients suffering from infection. Wright and Douglas did this in a large series of studies. In the case of staphylococcus infections the following experiment will illustrate their results:

TABLE I

(Wright and Douglas, *Proc. Royal Soc.*, Vol. 74, 1904.)

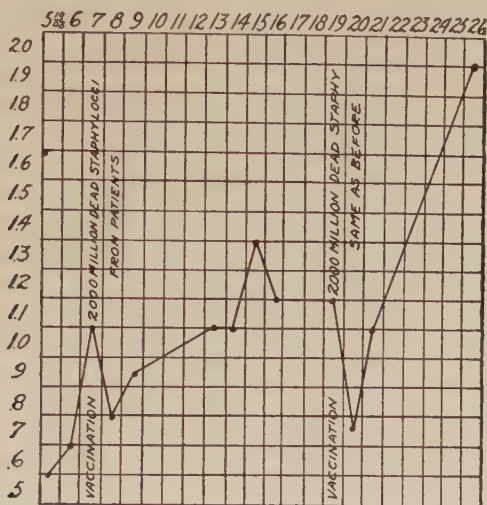
Showing the ratio in which the phagocytic or opsonic power of the patient's blood stood in each case to the phagocytic or opsonic power of the normal individual who furnished the control blood. (The phagocytic power of the control blood is taken in each case as unity.)

| Initials of Patient | Form of Staphylococcus Invasion | Opsonic Index |
|---------------------|---------------------------------|---------------|
| E. E. | Furunculosis..... | 0.48 |
| F. F. | Sycosis..... | 0.49 |
| J. E. | Acne..... | 0.64 |
| J. H. | Furunculosis..... | 0.87 |
| W. B. | Acne..... | 0.55 |
| E. H. | Acne..... | 0.82 |
| W. H. | Furunculosis..... | 0.79 |
| R. G. | Furunculosis..... | 0.7 |
| G. L. | Acne and sycosis..... | 0.74 |
| S. C. | Furunculosis..... | 0.87 |
| W. L. | Furunculosis..... | 0.88 |
| W. P. | Furunculosis..... | 0.39 |
| S. F. | Very aggravated sycosis..... | 0.1 |
| E. F. D. | Acne..... | 0.73 |
| D. C. | Sycosis..... | 0.8 |
| J. M. | Acne..... | 0.48 |
| W. M. | Sycosis..... | 0.37 |
| E. P. | Acne..... | 0.6 |
| M. S. | Pustular affection of lips..... | 0.6 |
| F. V. | Repeated staph. infection..... | 0.47 |

In this series, as in others investigated by Wright and his collaborators, staphylococcus infection was uniformly associated with a low index. He concludes that there is probably a causative relation between the two facts, in that under conditions of depressed phagocytic powers staphylococci may gain a foothold, while under

ordinary normal conditions they would fall prey to phagocytic destruction soon after entering the body.

The study of the opsonic index during the treatment of such cases with dead staphylococcus cultures (usually with the organisms cultivated from the patient's own lesions—"autogenous vaccines") revealed a striking coincidence between the rise of the opsonic index and improvement in the clinical conditions. A number of further interesting and practically important points were brought out by the systematic study of these relations which may be illustrated by



CURVE I.—RESULT UPON OPSONIC INDEX OF VACCINE TREATMENT IN TWO CASES OF CHRONIC STAPHYLOCOCCUS FURUNCULOSIS.

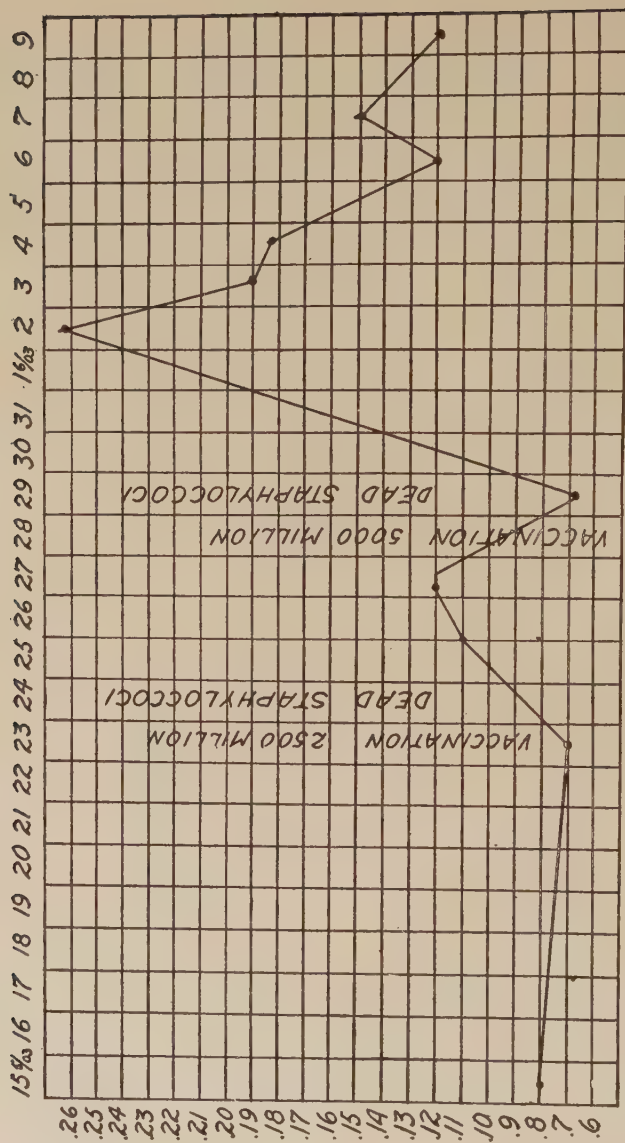
(After Wright and Douglas, *Proc. Royal Soc.*, Vol. 74, 1904, p. 156; also from "Studies on Immunity," p. 41.)

reproducing a plan of the opsonic index curves constructed from cases.

The curve shown above, and taken from a paper by Wright and Douglas, illustrates the course of the opsonic fluctuations in the case of a medical student who had suffered for four years from boils.

When first seen the opsonic index (1. being normal) was 0.6, and there were 2 boils on the neck. For 3 days after this there was a spontaneous rise in the index accompanied by an improvement of the lesions.

On the third day 2 billion staphylococci were injected. This was followed by an immediate drop of the phagocytic power—(the negative phase); together with this a new boil began to form. Soon, however, the opsonic power began again to rise, this time considerably above normal, reaching its highest point on the 8th day, when

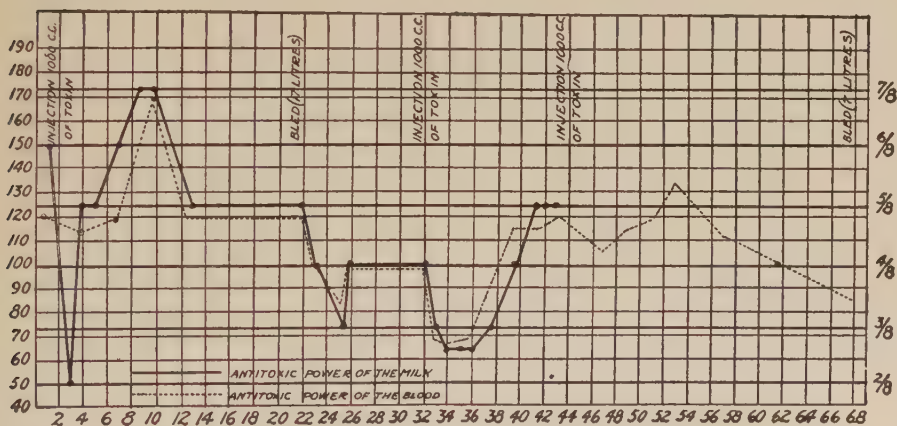


CURVE II.—THE RESULT OF VACCINE TREATMENT IN A CASE OF SEVERE SYPHILIS, TREATED BY WRIGHT AND DOUGLAS.
(After Wright and Douglas, *Proc. Royal Soc.*, Vol. 74, 1904, p. 156; also from "Studies on Immunity," p. 42.)

it again began to diminish. A second inoculation on the 12th day was followed by a similar preliminary negative phase, then a steady and rapid positive phase, which was accompanied by cure.

Another curve—Curve 2 of the same publication (Wright and Douglas, *Proc. Royal Soc.*, Vol. 74, 1904, p. 156)—is similar. This case suffered from severe sycosis (barber's itch), had been ill for 17 months, and had been unsuccessfully treated during this time with antiseptics. Staphylococci were isolated from a hair follicle, and from this the vaccine was made which was used in the treatment. Here the originally low opsonic index (0.8) rose after the first injection without a preliminary negative phase—but after the second treatment a sharp fall preceded the subsequent rise. Finally a sustained high index accompanied complete cure.

The rise and fall of the opsonins after the injection of bacteria is entirely analogous to the similar fluctuations of other antibodies after antigen injections. Measurements of this kind are numerous in the literature. Thus Salomonsen and Madsen, measuring the antitoxin contents of the blood and milk of a mare which were being immunized by injections of diphtheria toxin, obtained the following curve, which is entirely similar in essential features to those constructed for the opsonic index by Wright and Douglas:



CURVE DESCRIBING QUANTITATIVE MEASUREMENTS OF ANTITOXIN IN A MARE IN RESPONSE TO TOXIN INJECTIONS.

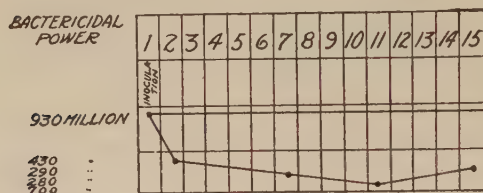
(Taken from article by Salomonsen and Madsen, *Ann. de l'Inst. Pasteur*, Vol. 11, 1897, p. 319.)

Results having the same general significance are apparent in the measurements made upon a tetanus toxin goat by Ehrlich and Brieger,⁹ and in the observations upon the fluctuations of bacteri-

⁹ Ehrlich and Brieger. *Zeitschr. f. Hyg.*, Vol. 13.

cidal power of the sera of patients treated with typhoid vaccines made by Wright¹⁰ himself. Similar, again, are the various agglutinin curves constructed by Jorgensen and Madsen¹¹ and others.

Apart from the purely theoretical value of such measurements, they demonstrate features which are therapeutically of the greatest importance. They show that in all processes of active immunization the injection of antigen is followed almost immediately by a rapid decline of specific antibodies in the blood serum. This "negative" phase, as it is called, is probably due to a neutralization of existing



PROLONGATION OF THE NEGATIVE PHASE DUE TO TOO VIGOROUS TREATMENT WITH TYPHOID VACCINE.

(After A. E. Wright, *Brit. Med. Journ.*, May 9, 1903. Also from "Studies on Immunity," p. 179.)

antibodies and lasts for varying periods, which must, of course, depend upon complex relations between the degree of resistance (or amount of antibody constituents of the serum), the quantity of antigen injected, and the general recuperative powers of the subject. Therefore, without some control like that furnished by the measurement of opsonins

or other antibodies it is impossible to determine whether the negative phase has ended or is still in progress unless the clinical condition is of such a nature or location that degrees of improvement or exacerbation are well marked and easily observed. Even then clinical observation alone is at best not an absolutely reliable guide.

The practical importance of the question lies in the harm which may accrue to the patient if a second injection is practiced before the cessation of the negative phase. Wright himself accentuates this danger by expressing the opinion that, in typhoid inoculations, an excessive dose administered to a patient in the physiological condition of the negative phase may be followed by a prolongation of this phase into a period of several months.

In the case of successive inoculations, as in vaccine treatment, a too rapid repetition—i. e., a repetition of injection during such a period of depression—leads to what Wright speaks of as a "summation of the negative phase," which obviously may seriously aggravate the condition of the case.

It is to such a cumulation of the negative phase that Wright attributes the failures attendant upon the use of tuberculin during the early days after its introduction, since injections at this time

¹⁰ Wright. *Practitioner*, Vol. 72, 1904, p. 118.

¹¹ Jorgensen and Madsen. *Festschrift*. Serum Institut. Copenhagen, 1902.

were carried out without any control of serum reactions in the patient and with comparatively large doses.

The danger to be carefully avoided, therefore, is a too rapid succession of inoculations and too large a dosage, since both of these procedures may be followed by cumulation of the "ebb tide of immunity," and great harm may result. On the other hand, if the treatment is so spaced and measured that the successive inoculations are given just before the positive phase has ended—in other words, just before the apex of the curve is reached—a moderate negative phase may be then followed by a second positive phase still higher than the first, and corresponding improvement will result. It is even possible to occasionally obtain a summation or cumulation of the positive phase—in which the negative phase will be entirely suppressed. This is illustrated in the following curve, in the case of the first and second inoculation indicated on the chart. This case, too, was a staphylococcus infection occurring in a laboratory attendant:



STAPHYLOCOCCUS INDEX AS DETERMINED BY WRIGHT IN A CASE OF ACNE TREATED WITH STAPHYLOCOCCUS VACCINES.

Note summation of positive phase after third injection. (After A. E. Wright, "Studies on Immunity," p. 348.)

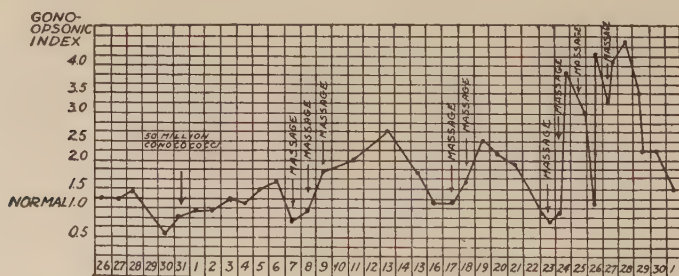
Such a summation of positive phase, though of course the ideal to be aimed at, cannot be produced with regularity, however carefully we may attempt to control the treatment. It is worth mentioning, moreover, a fact which should become evident from the preceding and is too often overlooked, that a summation of the negative phase can certainly be attained by the frequent repetition of larger doses. This is practiced not infrequently in the false hope of hastening the acquisition of immunity, and does harm more often than good.

Ordinarily the opsonic index when raised to a level considerably above normal will gradually recede to the normal or even to a sub-

normal condition. In isolated cases, however, especially in tuberculosis, the index may remain high for periods as long as a month. This Wright speaks of as a sustained "high tide" of immunity. These laws of fluctuation are all of them entirely analogous to those long well known in the cases of other antibodies, for even in diseases in which the immunity following an attack—(typhoid fever, cholera, plague, and others)—is continued through life the antibodies disappear from the blood after varying periods and we are forced to seek the cause of the permanently high resistance, not in the circulating blood, but in the ultimate physiological units—the cells and tissues.

According to Wright also, the treatment with vaccines may be either reënforced or entirely replaced by a process of autoinoculation from the patient's own lesion by increasing the local circulation, thereby throwing more of the specific antigen into the blood stream.

This reasoning has been applied, not only to the treatment of tuberculosis and other conditions, but has been utilized to explain fluctuations in the opsonic indices of untreated patients under the influence of unusual motion of the diseased parts—as in walking or other exercise. Wright's meaning is well illustrated by the following curve of opsonins in a case of gonorrheal polyarthritis in which massage of the joints resulted in reactions similar to those ordinarily elicited by vaccine injections:



OPSONIC CURVE IN A CASE OF GONORRHEAL ARTHRITIS IN WHICH AUTO-INOCULATION BY MASSAGE WAS PRACTICED.

(After Wright, Douglas, Freeman, Wells and Fleming, "Studies on Immunity," p. 373.)

A further modification of the vaccine treatment of Wright originated in the observation that the exudate present in many infected foci is often very much less rich in opsonins than is the blood serum of the same patient. This is not unlikely to be due to an absorption of the antibodies by the bacteria—as well as by the tissue detritus in the lesion. But Wright has interpreted it as a purely specific ab-

sorption by the bacteria, and has utilized it for diagnostic purposes. Thus, with Reid,¹² he has examined in this way the comparative amounts of tubercle bacillus-opsonins in the blood, and in the local exudates (peritoneal fluid) in cases suspected of tuberculosis, and has determined the tuberculous nature of the condition by showing a discrepancy between the two. These results have not been universally confirmed.¹³ But therapeutically, because of this supposed lack of opsonin in the fluid of lesions, Wright has advised the increase of the local flow of lymph by poulticing, heat, drainage, Bier's cups, X-rays, Finsen light, and other means of accomplishing this purpose.

All that has gone before (most of it taken directly from the staphylococcus studies of Wright and his immediate followers) has tended to show a very close correspondence of clinical improvement with the increased opsonin contents of the blood.

As applied to other infections, such as gonococcus arthritis, colon bacillus cystitis, localized pneumococcus lesions, and many other conditions of a localized character, observations of a similar general significance have been made. Such reports have been made, apart from the Wright school, by Emery,¹⁴ Potter, Ditman, and Bradley,¹⁵ Potter,¹⁶ Tunncliffe,¹⁷ Whitfield,¹⁸ Cole and Meakins,¹⁹ and many others, and we may say with reasonable accuracy that, in localized infections particularly, there is much evidence to show that clinical improvement and rise of the opsonic index go hand in hand.

There have been many exceptions to this—which, in view of the complicated factors involved in immunization, as well as the difficulty of the technique, is not surprising.

In tuberculosis—in which many of Wright's earlier studies were made—the parallelism has not been so consistent. Thus even the early work of Bullock²⁰ showed that, in contrast to similar staphylococcus investigations, the tuberculo-opsonic indices of patients may occasionally be higher than normal, and similar observations were made by Lawson and Stewart²¹ in cases of acute pulmonary tuberculosis.

Various investigations, too numerous to be reviewed in detail in

¹² Wright and Reid. *Lancet*, 1906; *Proc. Royal Society*, Vol. 77, 1906.

¹³ Opie. *Assoc. of Am. Phys.*, Washington, 1907.

¹⁴ Emery. "Immunity, etc.," Lewis, London, 1909.

¹⁵ Potter, Ditman, and Bradley. *Journ. A. M. A.*, Vol. 47, 1906, p. 1722.

¹⁶ Potter. *Jour. of A. M. A.*, Vol. 49, 1907, p. 1815.

¹⁷ Tunncliffe. *Jour. of Int. Dis.*, Vols. 4 and 5, 1907 and 1908.

¹⁸ Whitfield. *Practitioner*, May, 1908.

¹⁹ Cole and Meakins. *Johns Hop. Hosp. Bull.*, Vol. 18, 1907.

²⁰ Bullock. *Transact. of Lond. Path. Soc.*, Vol. 56, 1905, and *Lancet*, 1905, Vol. II, p. 1603.

²¹ Lawson and Stewart. *Lancet*, 1905, Vol. II, p. 1406.

this place, indicate in a general way that localized tuberculosis of the skin, joints, intestines, and glands, with the patient quiet and at rest, is apt to show a low index, while a high index may, under such conditions, often point to an active pulmonary lesion. According to Wright, this depends upon the following factors: In a localized lesion, with the body at rest—and when systemic symptoms such as fever are absent—the focus is, very probably, quiescent and in but slight communication with the circulation, even though it may be slowly progressive. In such cases little or no antigen is being discharged and, in consequence, no antibody formation is stimulated. Indeed, even the small amount of antibody which is present comes into but indifferent contact with the lesion because of its comparative insulation from the body fluids. Such a lesion may be benefited by rest, in that spreading is inhibited, and autointoxication, with the production of a negative phase, prevented; but it cannot be completely cured unless the antibodies are increased. This can be accomplished by carefully controlled vaccinations with tuberculin. At the same time more effective contact of these antibodies with the lesion may be attained by local applications, X-ray, etc. Or, again, the same purpose may be accomplished by carefully controlled and graded motion or massage of the diseased part—which may be used both to increase the opsonin contents by auto-inoculation and to enhance the local circulation. If this is done with care it may serve to substitute entirely for the treatment with vaccines.

On the other hand, such treatment with auto-inoculation, it must be remembered, is entirely uncontrollable as to dosage, and, therefore, not to be generally recommended.²² In active pulmonary tuberculosis, when there are systemic symptoms such as rise of temperature, the body is very probably already receiving excessive amounts of antigen and vaccine treatment of any kind may be dangerous.

However we analyze the work done on tuberculo-opsonins—and the investigations on this subject are far too numerous to be here reviewed—we are forced to the conclusion that in this disease the opsonic fluctuations are far more irregular than in most other conditions. Much,²³ for instance, found no regular differences between the tubercle bacillus opsonins of healthy and of diseased individuals, and Koehlich²⁴ obtained similar results, adding the important observation that animals that show a high natural resistance to the human type of the tubercle bacillus invariably show an opsonic index much lower than that of man.

We may question with much justice, therefore, whether in the

²² Meakin and Wheeler. *Br. Med. Jour.*, 2, 1905.

²³ Much. *Münch. med. Woch.*, p. 496, 1908.

²⁴ Koehlich. *Zeitschr. f. Hyg.*, Vol. 68, 1911.

case of this bacillus opsonic investigations can be looked upon as indicators of immunity with as much confidence as in cases of other bacterial invasions. It is true, indeed, that tubercle bacilli—as well as leprosy, rat leprosy, and other acid-fast bacteria—are eagerly taken up by polynuclear leukocytes when they are injected into the peritoneal cavity of a guinea pig or rat or other experimental animal. On the other hand, we have much evidence which seems to show that such phagocytosis is not in these cases a direct method of bacterial destruction. In another place we have cited the experiments of Tschernorutski,²⁵ which showed that polynuclear leukocytes, though containing other ferments, were devoid of lipase. And Carey and the writer—experimenting with rat leprosy bacilli—found that these acid-fast bacteria were not disintegrated within leukocytes in the course of weeks, while they were often subject to rapid destruction in the presence of living spleen cells in plasma. Furthermore, in the discussion of the tuberculin tests we have reviewed evidence which points to the fact that in the reactions to tubercle bacilli we have probably to deal more particularly with sessile receptors on fixed tissue cells than with specific circulating antibodies. Bartel and Neumann²⁶ have concluded that the phagocyte which takes up tubercle bacilli represents only a preliminary vehicle by which the micro-organisms are conveyed to the spleen and lymphatic tissues, in which actual destruction then takes place. While no final conclusions can be drawn from the available evidence, all these data render it uncertain whether the opsonic index as determined for polynuclear phagocytosis may be at all regarded as a reliable indication of increased or diminished resistance, and on this basis the control of therapy in tuberculosis by opsonin estimations is of course placed upon an uncertain basis.

We have then very briefly traced the work done upon opsonin determinations from the purely practical point of view. There is of course no question about the scientific accuracy of the observations upon which rests our knowledge of the opsonic properties of blood serum. There is also no doubt concerning our ability to increase the immunity of an individual by systematic treatment with vaccines made of pure cultures of bacteria. However, the work of Wright has concerned itself with two distinct questions which must be separately answered. Briefly stated these are: 1. What is the value of opsonic estimations in controlling the therapeutic vaccinations of patients? 2. To what degree and in which particular conditions may the process of vaccination (active immunization) be regarded as a hopeful method of therapy?

²⁵ Tschernorutski. *Hoppe-Seyler's Zeitschr. f. Phys. Chem.*, Vol. 75, 1911.

²⁶ Bartel and Neumann. *Wien. kl. Woch.*, Nos. 43 and 44, 1907; *Centralbl. f. Bakt.*, Vol. 48, 1909.

The first question has, in part, been answered in the preceding paragraphs. Reasonably accurate comparative estimations of the opsonic properties of serum can unquestionably be made by Wright's method, or some of its accepted modifications, in the hands of trained workers who look upon each estimation as an experimental problem and have time for control and repetition. That even in such cases the matter is difficult is amply testified to by such reports as that of E. C. Hort,²⁷ who states that two of the most skilled experts²⁸ in London, working with samples of the same serum taken before and after vaccination, reported—"the one that the index was raised, the other that it was lowered by the treatment." This, and similar experiments of other observers, do not, of course, invalidate the results obtained in special researches like those of Wright, Neufeld, and others, but they *do* indicate that the control of clinical cases by opsonic estimations is not a matter that can profitably be made a routine procedure by which the treatment of the cases can be regulated. As a problem of clinical research in a given series of patients opsonin studies are unquestionably valuable and the comparative data so obtained have proved, and will continue to prove, of great value. But we cannot hope as yet, it seems to us, to utilize this method, except in cases in which much time and care can be centered upon a few patients under the best conditions. Opinions essentially similar to this have been expressed by experienced clinicians (Potter,²⁹ for instance), who have followed out series of cases on which systematic opsonin determinations were made.

As to the opsonic index in tuberculosis, we believe that the experimental evidence at present available does not show that such measurements are reliable measures of resistance, and, in this disease, even when the index is taken with a degree of care which precludes gross error, it is doubtful whether its estimation is of as much value in controlling treatment as are the data obtained by skilled clinical observation.

This leaves us, therefore, for the control of vaccine treatment in the routine work of the clinic only the information gleaned from such indications as alterations in any visible or palpable lesions, general systemic symptoms, temperature, leukocytosis, etc. Since these will present such manifold and variable pictures in different conditions, generalization is useless.

The second question concerning the value of vaccine treatment in infectious disease of human beings cannot be so briefly answered, and is one of the greatest importance in medicine. It is well known

²⁷ Hort. *Br. Med. Jour.*, Feb., 1909, p. 400.

²⁸ Quoted from Adami, *Trans. Amer. Phys. & Surg.*, Vol. 8, 1910. See also Pearson, *Biometrika*, 1911.

²⁹ Potter. *Loc. cit.*

that tuberculin therapy has come into carefully controlled use in recent years only, although it was introduced early in the history of specific therapy by Koch. The misuse and failure of this treatment during the years following its introduction are easily explained by the defective knowledge of antibody reactions and the general principles of immunity—a condition which was removed only by the subsequent assiduous work of numerous investigators. At the present time the value of this method of treatment is being acknowledged, though its limitations and possible dangers are properly recognized. The Wright method of vaccine treatment is also an unquestionably powerful therapeutic weapon, and yet, owing to commercialization, unskilful application, and, more especially, because of extensive attempts to apply it in unsuitable cases, it may easily, like tuberculin therapy, enter into a period of neglect and disrepute. It is very necessary to accentuate at the present time that the active immunization of human beings with any form of bacterial product is a serious procedure which requires painstaking and skilled control, and should not be undertaken without the same degree of preliminary experience and study which is considered prerequisite in any other branch of specialized medicine.

Any opinion expressed regarding the ultimate value of a method of treatment which is still undergoing active clinical investigation must of course be purely tentative. Moreover, there are so many differences of judgment that we wish to emphasize the purely personal character of the views expressed.

In passing judgment upon the value of active immunization in man we must distinguish sharply between active immunization which is prophylactic and that which is carried out after the disease has gained a definite foothold in the body. In the former case we are dealing with a new method and with one upon which the very foundations of our knowledge of immunity have been built. It is the method of Jenner in small-pox. It is that of Pasteur in chicken cholera, in anthrax, and in many other infections. It has been used as a routine in animal experimentation in laboratories since the first days of the systematic study of infections. There is no question about its being a rational and logical procedure. The immunity which can be easily conferred upon a healthy individual in this way need not be extensively above the normal in order to protect from invasion by the small numbers of pathogenic germs which may gain entrance under conditions of accidental, spontaneous infection.

The possibilities of the method were recognized by Ferran, a pupil of Pasteur, who applied it to cholera, and, since his time, it has been extensively attempted in many of the infectious diseases which occur epidemically, and therefore justify attempts in this direction.

In essence also Pasteur's method of active immunization in rabies represents such prophylactic vaccination, since, in this case, although treatment is begun after infection has taken place, nevertheless the process of immunization is carried out during the incubation period before active manifestations of the disease have set in. Prophylactic vaccination, therefore, is a valuable procedure which has reaped remarkable results of recent years, especially in protection against typhoid fever. In a subsequent section this phase of vaccination is more extensively discussed, and we may therefore leave it for the present.

In this place we are more particularly concerned with the problem of the treatment of existing disease with vaccines prepared from the bacteria by which the disease is caused. In how far this is justifiable or even logical is a question which depends upon the conditions of each individual case. We can approach the problem best by roughly classifying the various forms in which infection occurs in the human being.

When bacteria gain entrance into the tissues of the human body, granted that the organisms are pathogenic, an immediate struggle ensues between the offensive properties of the micro-organisms and the defensive powers of the tissues. The factors which determine the outcome of such a combat have been more fully considered in Chapter I. Briefly, if the defensive powers of the body greatly preponderate the result is localization and rapid destruction of the micro-organisms—with cure. In such a case any form of treatment is unnecessary. On the other hand, the balance of power may be turned in the opposite direction, in which case the infectious process becomes rapidly generalized, the bacteria enter the blood stream and lymphatics, and the defensive powers are overwhelmed. In such a case also active immunization with vaccines is entirely useless.

There are cases, however, in which the struggle is a more equal one, and in which the infectious process is held in check by the defenses, so that it takes a slow, chronic, localized form, and spreads, if at all, very slowly. What is it in such a case that prevents complete healing of the process? The answer to this may be found both in local and in systemic causes. Locally the lesion, after the preliminary skirmishes, may become encapsulated either by fibrin formation, clot, or other tissue changes so that, as Wright suggests, the fluid constituents of the blood-plasma cannot easily approach the organisms in the lesion. The same effect may result from internal pressure by fluid and possibly by the presence of considerable quantities of tissue detritus, by which protective serum constituents are fixed and thus diverted from the bacteria. Against these factors, of course, no form of immunization can be of value. Wright recognizes this, and suggests the use of surgical evacuation, Bier's method,

X-rays, Finsen light, heat, and a number of other localized methods of increasing the blood supply. This, too, may be the reason for the benefits derived from wet dressings, in that they keep the tissues macerated, soft, and moist. At any rate, it is a matter of local surgical treatment. At the same time, however, there may be systemic causes which prevent the complete healing of such lesions, namely, an insufficient supply of circulating antibodies, opsonic or bactericidal substances. These may be sufficient to hold the lesion in check, but since small quantities of bacteria only are in contact with the blood stream, relatively small amounts of antigen are absorbed and antibody formation is consequently deficient. Here we have an ideal condition for vaccine therapy. By isolation of the organisms from the patient's lesion, for which, in this case, there is time, and the careful immunization of the patient with these organisms, the immunity may be considerably increased and cure effected.

Closely related to this type of lesion are those conditions in which there are localized infections which heal rapidly but recur in quick succession again and again. Such are the common cases of consecutive crops of boils; and not dissimilar are the manifestations of erysipelas where the lesion extends along the edges while it heals in the center. There is in this type, probably, a very close balance between protection and offense; the defensive reaction is sufficient to overcome the localized lesion, but insufficient to set up a permanent systemic protection. A certain amount of local immunity acquired by the tissues of the affected areas may suffice to throw slight weight into the balance on the side of protection, enough at least to decide the struggle; and this element of locally acquired tissue resistance is in all probability also the cause for the failure of these lesions to recur immediately in the same area. Here, too, treatment with vaccines is not illogical and may yield good results if properly carried out.

In generalized systemic infections we must sharply distinguish between cases of acute sepsis in which the bacteria are actively growing and multiplying in the circulation and cases in which blood cultures are positive only because the bacteria are being constantly discharged into the circulation from a focus in the tissues. In the former the defenses of the body are overwhelmed by an extensive flooding with the bacteria, and vaccines, if not harmful, are, at any rate, utterly useless since the antigen is already so extensively distributed throughout the tissues that if the body were capable of responding with sufficient antibody formation this would unquestionably occur without the small additional amount furnished in the bacterial emulsion. Vaccination in such cases is entirely analogous to an attempt to stimulate a degenerated heart muscle with strychnin—the whipping of a tired mare.

Such cases of septicemia, however, are not in our opinion the most common ones in the human being. It is probable that all localized infections of more than a very trifling nature discharge living bacteria into the circulation from the very beginning. However, in most cases the bacteria, though able to hold their own in their entrenched position at the focus where accumulated offensive factors and local injury reënforce them, are yet rapidly destroyed when, in small detachments, they get into the open circulation where the plasma antibodies and phagocytes are freely active. There are cases which take a middle course between such purely localized lesions and the acute septicemia, conditions in which a well-established focus continues to furnish bacteria to the blood stream as fast as they are destroyed. An example which illustrates our meaning well is that of the so-called subacute endocarditis caused by the *Streptococcus viridans* and its close biological kin, where blood cultures are often consistently positive for a long period or may show occasional intervals in which the blood is bacteria-free. The focus on the heart valves apparently can continue uncured in spite of a relatively high or at least normal systemic resistance to the microorganisms. If, as we ourselves have done, we isolate the organisms by blood culture from such cases, and then measure the opsonic properties of the patient's own serum against them, using the patient's own leukocytes, we may often find that active phagocytosis takes place, in a degree equal or even superior to that taking place in the serum of normal individuals. Neither does there seem to be a diminished phagocytic power of the patient's own leukocytes. For a long time these conditions may continue, with a constant destruction of bacteria in the blood and a corresponding renewal of the supply from the lesion. The same condition can be observed in rabbits in which chronic endocarditis with persistently positive blood culture has been produced by injections of these bacteria. In such animals measurements similar to those described above have been made by Miss Gilbert in our laboratory, and it has seemed as though persistently positive blood cultures could be obtained only when a localized focus was set up in the animals. Unless this is the case the blood cultures rapidly become negative.

Conditions essentially similar may exist in any other form of severe localized infection. Positive blood cultures do not necessarily mean a multiplication of the bacteria in the blood stream and a rapid overwhelming of the body. We have had occasion to see a number of cases of bacteriemia in which the focus of infection was surgically accessible; and in some of these cases early removal of the focus and purely surgical treatment resulted in a clearing up of the infection. Similar experiences have been reported by Libman and a number of others, and for this reason general septicemia, if not fulminating, may still be less desperate than ordinarily supposed.

Now, having outlined the conditions obtaining in such cases, let us briefly consider whether, under the circumstances, vaccine therapy may logically be regarded as a hopeful form of treatment. We may assume, on the one hand, that the bacteria, being consistently present and destroyed in the blood, should furnish antigen sufficient to stimulate the body tissues to their utmost reactive ability. This would seem a strong argument against vaccine therapy. On the other hand, we must take into consideration another phase of the subject, one which has some experimental justification. In discussing the origin of antibodies in another section it will be remembered that we called attention to the fact that many different tissue cells probably participate in the production of these protective reaction-bodies. We cited an experiment of Wassermann and his pupils in which they proved that antibodies were produced most energetically in the tissues about the point of injection of the antigen, namely, in the place at which it came into most concentrated contact with the cells. They injected bacteria into the subcutaneous tissues of the ear of a rabbit, measured the progressively increasing appearance of antibodies in the blood stream, and then amputated the ear. A sudden drop of antibody contents followed, showing that the supply of antibodies had largely emanated from the tissues surrounding the injection point. Park³⁰ has pointed out another reason why vaccine treatment may be expected to exert beneficial action in such cases. He calls attention to the fact that when very large amounts of antitoxin are added to toxin before injection no antibody production results, and assumes that in chronic or subacute general infections the circulating bacteria are in contact with specific antibodies, partially "sensitized," and therefore not efficient as antigen. In consequence the injection of homologous unsensitized bacteria may hasten antibody formation. This assumption of Park is theoretically valid, but it is not in accord with the more recent experiments of Metchnikoff and Besredka, who claim to have obtained the best results in prophylactic typhoid vaccination by the injection of sensitized bacteria.

Thus the use of vaccines in the subacute or chronic cases of infection with bacteria in the blood stream may be theoretically justified, and no one can say at the present time whether or not it has therapeutic promise. At any rate, it cannot be absolutely condemned on theoretical grounds.

Like so many other phases of this question, it must be answered ultimately by clinical experience, for in experimentation upon animals, while it is easy to produce a purely localized lesion followed by rapid healing, or a generalized lesion leading to rapid death, it is not easy to produce prolonged infections with anything like regularity, and there are so many modifying accidental factors which

³⁰ Park. *Trans. of Amer. Phys.*, Vol. 8, 1910.

influence the course of such infections in animals that the results of vaccine treatment in them are difficult to judge.

In acute diseases which run a definite course, typhoid fever, pneumonia, dysentery, cholera, plague, and a number of other conditions, vaccine treatment during the course of the disease has not much justification. In typhoid fever, especially, specific antibodies appear in the blood in amounts enormously increased above the normal at periods when the patient is still actively ill in spite of the fact that the blood stream has been freed of the micro-organisms. Whatever may be our opinion as to the continuance of the disease after bacteria have been driven out of the blood stream, the use of vaccines can only tend to further increase of antibodies which are already present in amounts far exceeding normal. In pneumonia the micro-organisms seem curiously resistant against the attack of the serum antibodies, and in spite of the presence of large amounts of antigen both in the lungs and, for a time, in the circulation the development of immunity is delayed until just before or near the crisis. Since this, however, is usually only a matter of 7 or 8 days, it is hardly likely that the injection of vaccines during this period could markedly alter the ultimate outcome. In plague we have usually an acute septicemia, and here the considerations that we have outlined above are applicable.

There are none of the acute infectious diseases of specific course in which vaccine treatment after onset seems advisable on theoretic grounds.³¹

As we have stated before, the opinions expressed above are given with the purpose of stating as clearly as we can the logic of vaccine³² therapy as we see it at present. The next ten years of clinical experience may largely modify these views. One thing is certain, however, and that is that the problem can only be settled if treatment by this method is undertaken with the guidance of an accurate bacteriological diagnosis, and with bacteriological control of the individual case, so that, when occasion arises, estimations of antibodies can be made.

To protest against the random use of commercial stock vaccines without laboratory diagnosis and without control is almost a platitude.

In the case of tuberculosis the problem had been actively investigated before Wright, and there seems little question that tuberculin therapy properly and cautiously applied has an established value in the treatment of initial and localized tuberculous disease. Whether

³¹ See also Theobald Smith, *Jour. A. M. A.*, Vol. 60, 1913, and R. M. Pearce, *Jour. A. M. A.*, Vol. 61, 1913.

³² For discussion of various clinical applications of vaccine treatment see symposium on vaccine treatment, *Trans. of Ass'n of Amer. Phys. and Surg.*, Vol. 8, 1910.

or not its use in actively progressive tuberculosis may or may not be hopeful, in which particular cases, and by what methods, it is to be applied, these are problems that we have neither the space to deal with nor the experience to summarize properly. They constitute a special field of clinical research, a survey of which may be obtained in such works as that of Bandelier and Roepke,³³ or the more especial experimental studies of Denys.³⁴

THE PRODUCTION AND STANDARDIZATION OF VACCINES

Vaccines in the sense of Wright consist merely of killed cultures of the bacteria with which the patient is infected. In all cases it is extremely desirable to make such vaccines "autogenous," by which we mean that the organism used is one which has been isolated from the case. The difference between various strains of the same species of bacteria seems to make this imperative whenever it is at all possible. The recent investigations of Neufeld and Haendel in determining that there are a number of types of pneumococcus which are antigenically distinct illustrates this point. The same principle is made clear by the recent work of Rosenau on the streptococcus-pneumococcus group. Especially important is Rosenau's observation that a pneumococcus which he had been able to transform culturally by special methods was found to be altered also in its reaction to agglutinins.

In the development of prophylactic methods of vaccination against epidemic disease like typhoid, cholera, plague, etc., many different methods of antigen preparation have been developed. In typhoid prophylaxis the bacteria have been used dead, living, and sensitized, and even extracts have been employed. In cholera the early use of living cultures by Ferran has given way, in the hands of Kolle and others, to that of dead bacterial emulsions. In plague and a number of other conditions the impression seems to be general that the bacteria should be used in the living, but attenuated, state. Special methods which have been developed in these cases are discussed in another section.

In treatment of developed diseases with vaccines the method most commonly used is that which has been introduced by Wright, namely, the use of dead cultures. In his earlier experiments Wright cultivated the bacteria on agar slants for about 24 hours, then washed off the growth with 10 c. c. of sterile salt solution. It will be well to describe in detail the preparation of such a vaccine.

The bacteria must be isolated from the patient by the usual method of plate cultivation and colony fishing on suitable media.

³³ Bandelier and Roepke. "Lehrbuch der spez. Diagnostik und Therapie der Tuberkulose," 6th Ed., Kabitzsch, Würzburg, 1911.

³⁴ Denys. "Le Bouillon Filtré," Louvain, 1903.

We do not think that any satisfactory substitute for careful isolation by plating has been devised. After a pure culture of the organism has been obtained this is grown on relatively large surfaces of agar, glucose-agar, or ascitic agar, as the case may require. These cultivations may be made in Kolle flasks or, as Wright and others have suggested, on large agar surfaces obtained when the culture medium is allowed to harden in a square 3-oz. medicine bottle laid on its side. Any device of this kind in which a large surface of agar is exposed may be used.



CAPSULE MADE
OF TEST TUBE
TO HOLD STOCK
VACCINE
EMULSION
FROM WHICH
DILUTIONS
ARE MADE.

(It is well to keep capsule open to capillary tip until it has cooled off, otherwise it may crack when quickly cooled.)

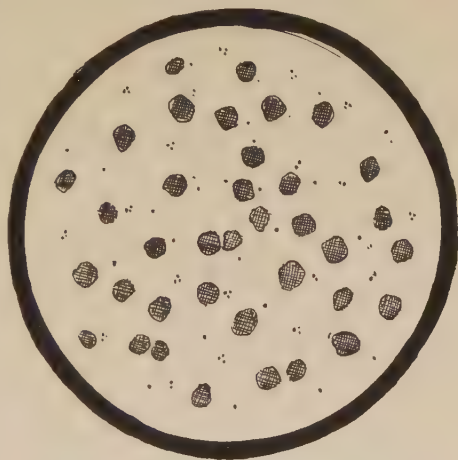
After suitable growth of the micro-organisms has taken place, 24-48 hours, the growth is gently washed off with 10 c. c. or more of sterilized salt solution. Care must be taken to do this in such a way that no agar is drawn away with the emulsion. The thick emulsion so obtained is removed from the culture bottle with sterile nipple pipettes or Pasteur pipettes and transferred to a sterile thick-walled test tube into which glass beads have been placed. By drawing out the neck of this test tube in the flame a glass capsule is formed, in which we now have our so-called stock emulsion. (See figure.)

The next thing to be done is to standardize this stock emulsion, or, in other words, determine approximately the number of bacteria to the cubic centimeter. There are a number of methods by which this can be done.

The method most extensively used by Wright and his followers was that in which the bacteria are counted against red blood cells. The bacteria in the capsule are shaken thoroughly with glass beads so that clumps may be broken up and even distribution obtained. A little of the emulsion is then put into a clean watch glass, a step which can be accomplished most easily by breaking off the tip of the drawn-out part of the capsule, tilting it very gently and heating the closed end over a small flame, so that some of the emulsion will be driven out by the expanding air. With a nipple pipette marked about an inch from the tip, as in the taking of an opsonic index, a little of the emulsion is drawn up. This is placed into another clean watch glass and is mixed with about 2 volumes of salt solution and one volume of blood from the finger, these quantities being measured with the same nipple pipette. We then have a mixture in which, in a total of 4 volumes, there are equal parts of blood and of bacterial emulsion. After this emulsion has been thoroughly mixed by drawing in and out through the nipple pipette smears are made on

slides and stained with Jenner or any other suitable blood and bacterial stain. Under the field of the microscope the ratio between the bacteria and blood cells is then determined, and from our knowledge of the number of the red blood cells in this blood to each c. mm. we can easily calculate the number of bacteria to the c. mm. or c. c.³⁵

A more accurate method of enumerating the bacteria in a suspension to be used for vaccine is by direct count of an accurately made dilution in a hemocytometer chamber, as was first suggested by Malory and Wright in 1908.³⁶ The bacterial suspension is diluted in blood-counting pipettes, 1-20 to 1-100 dilutions of thick bacterial suspensions being as a rule satisfactory. As a diluent one may use either salt solution or some dilute anilin dye, such as one made by mixing one part alcoholic methylene blue with 40 parts of 1 per cent. carbolic acid. The dilute suspension is then placed in an ordinary Thoma-Zeiss chamber, which was designed for counting blood platelets and has a depth of 0.02 mm. This enables one to use an oil immersion lens or high power dry system with a short working distance. From such a count one may readily estimate the number of bacteria in the original suspension; for example, if 20 squares in the Helber-Zeiss chamber are counted the result gives the number of bacteria in 0.001 c. mm.³⁷



MICROSCOPIC FIELD AS SEEN IN STANDARDIZATION OF VACCINES BY WRIGHT'S METHOD.

Another method of standardization of vaccines which is sufficiently accurate for clinical purposes is that of Hopkins, which consists in measuring the volume of the sediment³⁸ after centrifugalizing the preparation under standard conditions in a graduated tube. The tubes may be made with a capacity of 10 to 15 c. c. with a capil-

³⁵ For such counts it is convenient to contract the field of the microscope by using a diaphragm or simply marking a circle on the eyepiece with a grease pencil.

³⁶ Malory and Wright. "Pathological Technique," 4th Ed., New York, 1908.

³⁷ Glynn, Powell, Rees, and Cox. *Jour. of Path. and Bact.*, Vol. 18, 1914, p. 379.

³⁸ Hopkins. *Jour. A. M. A.*, 1913, Vol. 60, p. 1615.

lary tip about one inch in length, having a capacity of about 0.05 c. c. graduated in 0.01 c. c. The bacterial suspension, after being filtered through sterile cotton to remove fragments of the agar or other foreign bodies, is centrifugalized in such a tube for half an hour at about 2,800 revolutions a minute. The supernatant fluid and bacteria are removed down to the 0.5 c. c. mark and the sediment resuspended in 5 c. c. sterile salt solution by means of a capillary pipette which gives a 1 per cent. suspension. 0.05 c. c. of streptococci sedimented in this way represent quite constantly 16 mm. of dried bacterial substance. The number of organisms per cubic centimeter contained in 1 per cent. suspension in this way are as follows:

| | |
|-------------------------------------|------------|
| Streptococcus aureus and albus..... | 10 billion |
| Streptococcus..... | 8 " |
| Gonococcus..... | 8 " |
| Pneumococcus (capsulated)..... | 2.5 " |
| Bacillus typhosus..... | 8 " |
| Bacillus coli..... | 4 " |

After the vaccines have been standardized suitable dilutions can be made in salt solution to which 0.5 per cent. carbolic acid or some other antiseptic has been added. The dilutions are usually so made that from 100 to 500 million bacteria are contained in the cubic centimeter, this being a suitable initial dose of most organisms. The dilutions are placed in sterile bottles containing beads and fitted



VACCINE STOCK
EMULSION IN
RUBBER TOP
BOTTLE.

with rubber caps. These bottles can be shaken before use, the emulsion thoroughly distributed, and the desired quantity can be taken out with a sterile hypodermic syringe thrust through the rubber cap after this has been covered with a small amount of lysol or strong carbolic (see figure). After the dilutions have been made both these and the stock vaccines should be sterilized. Some workers sterilize always the stock vaccines and make the dilutions with aseptic proportions in such a way that no further sterilization is necessary. This is preferable because the less heat that is applied the better it is for the

preservation of their antigenic properties—sterilization is usually accomplished by heat in the water bath. Wassermann's earlier technique called for heating to 60° C. for one hour for a number of consecutive days. It is generally considered at the present time that it is better not to heat above 55° C. After the vaccine has been heated its sterility must be controlled to aerobic and anaerobic cultivation, and possibly by animal inoculation, although, except in special cases, this is unnecessary. Some workers,

especially when the vaccine is to be extensively used, as in typhoid immunization, inject some of the vaccine into white mice to exclude the possibility of contamination with tetanus. In such cases also it is not inadvisable to test out the antigenic value of the vaccine upon animals, measuring the agglutinins, etc., which result from a number of inoculations. In the preparation of a therapeutic vaccine where speed is required this of course is not feasible. Moreover, it is unnecessary in view of the fact that we wish to inject that particular organism into the patient from whom it has been cultivated. Whatever its antigenic value may be from animal experiments, it is preferable for the given purpose to any other strain.

Sensitized vaccines are easily made by exposing emulsions of the bacteria to moderate amounts of a strong immune serum which has been heated to 56° C. to destroy the complement. Bacteria will usually agglutinate under these circumstances and can easily be centrifugalized to the bottom. The excess serum is then washed off and the bacteria emulsified as in the case of the preparation of vaccines with dead organisms.

THE TUBERCULINS

Since we shall not attempt to discuss critically tuberculin treatment, as this is a subject upon which many special studies have been made both by clinicians and by laboratory workers, and is entirely too extensive to be reviewed in a book like this, on the other hand, we deem it a part of our task to discuss at least the methods by which the antigen or tuberculin preparations are obtained. There has been much discussion concerning the nature of the antigenic substances obtained from the tubercle bacillus. It has been claimed by Denys and others, for instance, that the tubercle bacillus may give rise to small quantities of a true exotoxin with consequent endotoxin-inducing properties. Again, most observers have believed that the poison of the tubercle bacillus consists of substances comparable to the endotoxin of other micro-organisms. The matter is by no means settled, and without going into the theoretical aspects of the problem we will confine ourselves in this place to a description of the production of the various forms of so-called "tuberculin."

OLD TUBERCULIN (KOCH)

The first tuberculin prepared by Koch is made in the following way: Tubercle bacilli of the human type are grown for from 4 to 6 weeks upon a 5 per cent. glycerin broth. The cultures are then sterilized in an Arnold sterilizer and are evaporated at about 80° C.

to one-tenth the original volume. This 60 per cent. glycerin extract of the tubercle bacilli is then filtered clear and constitutes the tuberculin.

The old tuberculin is a preparation which is extensively used in the subcutaneous and intracutaneous tests upon human beings and cattle, and forms the basis of the various preparations by von Pirquet, Moro, and others in the cutaneous tuberculin reactions. In his earliest work von Pirquet used a 25 per cent. solution of the old tuberculin. At present an undiluted old tuberculin is used for these purposes.

The old tuberculin also is the material from which the preparation for the ophthalmotuberculin test is made. For this purpose Calmette advises precipitating old tuberculin with double the volume of 95 per cent. alcohol, allowing the precipitate to settle and repeatedly washing the sediment with 70 per cent. alcohol. The powder which results is thoroughly dried, pulverized, and made up for use in 0.5 per cent. solutions. Bandelier and Roepke recommend the use of the diluted old tuberculin directly for these tests, employing a 1 per cent. solution.

NEW TUBERCULIN (T R AND T O)

The description of the preparation of these tuberculins we take from Ruppell in the *Lancet*, March 28, 1908. Virulent cultures of tubercle bacilli are dried in the vacuum and are then thoroughly pulverized by specially constructed machinery, and the grinding is continued until no intact bacilli are found in the preparation. One gram dry weight is then shaken up in 100 c. c. of sterile distilled water. The mixture is then centrifugalized at high speed—the supernatant fluid is T O (tuberkulin ober-schicht). This contains the water-soluble substances of the bacillus and gives no precipitate with glycerin. The residue—T R (tuberkulin ruckstand)—is again dried and ground up, shaken up in water, and centrifugalized. This is repeated 3 or 4 times, the total volume of water used for all the repetitions not exceeding 100 c. c. At the end of several repetitions all the T R goes into emulsion, and the various supernatant fluids obtained during these repeated grindings and shaking are mixed together and constitute the final T R preparation. This preparation, according to Koch, contains important antigenic substances, it gives a precipitate with glycerin, and it is standardized by the determination of the solid substances contained in a cubic centimeter. This, for a standard preparation, should be 0.002 gram to a cubic centimeter.

NEW TUBERCULIN BACILLARY EMULSION

This preparation consists of a combination of T O and T R. It represents an emulsion of pulverized tubercle bacilli in 100 parts of 50 per cent. glycerin. The preparation as marketed contains 0.005 gram solid substance to the cubic centimeter. It is prepared simply by mechanically grinding the bacteria as in the new tuberculin, but, instead of centrifugalizing for the separation of T O and T R, the bacteria are allowed to sediment after the addition of glycerin. This is the preparation which is extensively used in many places at present for the treatment of tuberculosis. It was adopted by Koch particularly because of experiments in which he showed that the treatment of animals with such preparations greatly increased the agglutinins for tubercle bacilli.

BOUILLON FILTRÉ (DENYS)

Denys cultivates the tubercle bacilli upon 5 per cent. glycerin bouillon as in the preparation of old tuberculin, but does not heat, sterilizing his cultures by filtration through porcelain. Denys believes that the application of heat in sterilization destroys exotoxins which have valuable antigenic properties.

SENSITIZED TUBERCULIN

Following the introduction of sensitized vaccines in other diseases by Besredka, Meyer³⁹ has introduced the sensitized tuberculin. This tuberculin is prepared in the following way: Tubercle bacilli of the human type are washed and dried and are mixed with a considerable quantity of the serum of animals immunized with tubercle emulsions and containing considerable quantities of tubercle-agglutinins. These serum mixtures are kept at 37° C. for several days and are then shaken in a shaking machine until intact tubercle bacilli are no longer to be found. The tubercle bacillus fragments are then thrown down in the centrifuge, washed in salt solution, and emulsified in 40 per cent. glycerin, 0.5 per cent. carboic acid being used. The emulsion contains 0.005 gram dry weight to a cubic centimeter. We take the description of the preparation from that cited by Bandelier and Roepke.

The above tabulation contains the most important tuberculin preparations as they are at the present time in use. For detailed studies of their clinical application we refer the reader to the very valuable book of Bandelier and Roepke, "Lehrbuch der spezifischen Diagnostik und Therapie der Tuberkulose," Curt Kabitzsch, Würzburg.

³⁹ Meyer. Cited from Bandelier and Roepke, "Lehrbuch d. spez. Diagn. u. Ther. d. Tuberkulose," Kabitzsch, Würzburg, 6th ed., 1911, p. 186.

CHAPTER XV

ANAPHYLAXIS

FUNDAMENTAL FACTS

THE fundamental principle of active immunization is the fact that the treatment of animals with bacteria or bacterial products, carried out according to certain empirically determined methods, leads to increased tolerance or resistance. The limitations within which this statement is true, and the variable factors to which it is subject, we have considered in the foregoing discussions dealing with the antibody-antigen reactions.

Although these reactions were studied at first purely from the point of view of increased resistance to infection, the most extensive studies of antibody formation have been made with such antigens as blood cells, serum, and other substances which are in themselves entirely harmless. For, in such reactions, great simplicity and ease of experimentation could be attained. For a time, therefore, the primary problem of increased tolerance or resistance was relegated to a secondary position, or, at least, dealt with chiefly by analogy, and the phenomena of increased antibody formation and increased resistance to the antigen were assumed to maintain a more or less strict parallelism.

That the problem is not as simple as this has gradually become obvious. We have come to recognize that the treatment of animals with any antigen, bacterial or otherwise, though leading to increased tolerance under certain conditions and within definite limits, may, under other conditions, give rise to the very opposite, that is, to an intolerance or increased susceptibility.

The development of this knowledge, like much else that serum study has revealed in the last fifteen years, takes root in isolated observations scattered throughout the early literature, but often regarded as merely noteworthy accidents or technical errors. This particular problem, moreover, was confused by the fact that some of the earliest observations regarding hypersusceptibility were made in the course of experimentation with diphtheria and tetanus toxins, antigenic substances toxic in themselves and, therefore, as we shall see, clouding some of the basic principles apparently involved in the phenomenon of which we now speak as anaphylaxis. We will for the present, therefore, limit our discussion to the development of the

knowledge of anaphylaxis merely as it concerns the hypersusceptibility incited in animals and man by treatment with various antigens, such as animal sera and other proteins, which possess but slight native toxicity or no toxicity whatever in themselves.

The special problem of toxin hypersusceptibility ("Giftüberempfindlichkeit" of von Behring) we will deal with later in a separate section, since it is as yet very doubtful whether these phenomena may justly be incorporated with true anaphylaxis as we now define it, despite the admitted fact that attention was called to the problems of acquired susceptibility largely because of these toxin investigations.

The earliest observation having direct bearing upon protein anaphylaxis is one which Morgenroth discovered in the writings of Magendie. Morgenroth¹ mentions that, in his "Vorlesungen über das Blut," published in 1839, Magendie describes the sudden death of dogs which had been repeatedly injected with egg albumen. Although Morgenroth, whose paper was written before the present facts regarding hypersusceptibility were fully developed, attributes these results to the action of precipitins, there can be little doubt as to the anaphylactic nature of Magendie's results.

A clear statement of the fundamental phenomena was given, also, by Flexner,² in 1894. In describing certain experiments he says: "Animals that had withstood one dose of dog serum would succumb to a second dose given after the lapse of some days or weeks, even when this dose was sublethal for a control animal."

One of the experiments cited to justify this statement is as follows:

"Two rabbits received $\frac{1}{2}$ of 1 per cent. and 1 per cent. of their body weight respectively of dog's serum, twenty-four hours old, on January 19, 1894. With the exception of hemoglobinuria, indisposition to move, and increased respiration, no ill effects were noted. The animals still showed hemoglobinuria on the following day. These symptoms disappeared and apparently the rabbits entirely recovered. On February 12, 1894, each received 1 per cent. of their body weight of dog's serum intravenously. A control animal also received 1 per cent. of its body weight of the same serum. The two animals that had been previously inoculated died in two and twelve hours respectively; the control animal showed only hemoglobinuria which disappeared after a day or two."

The experiment here quoted is, as a matter of fact, a perfect example of what we now know as "active sensitization."

However, the isolated observations recorded above were neither correlated nor followed out to their logical developments, and a

¹ Morgenroth. "Ehrlich Gesammelte Arbeiten," Transl., Wiley & Son, N. Y., 1906; p. 332 footnote.

² Flexner. *Medical News*, Vol. 65, p. 116, 1894.

systematic and purposeful study of the problem was deferred until Richet and Portier³ attacked it in 1902.

Richet and Héricourt⁴ had observed in 1898 that dogs treated with eel serum, which is toxic *per se*, could be killed by a second injection of an amount too small to injure normal untreated animals. Some years later Richet, in collaboration with Portier,⁵ determined a similar fact in the case of a poisonous substance, "actinocongestin," which they isolated by extraction of the tentacles of actinia.

Some of the facts of Richet and Héricourt's observations are as follows: Actinocongestin injected intravenously into dogs in quantities of 0.05 to 0.075 gram per kilo weight may cause illness, with vomiting, diarrhea, and respiratory distress, but does not kill. A dose of 0.002 gram per kilo causes no symptoms in a normal dog. If, however, 0.002 gram of the poison is injected into a dog which has previously received a sublethal dose and recovered, the result is violent illness and often death. It was obvious, and this was clearly stated by Richet, that the first dose had induced a condition of markedly greater susceptibility to the poison.

He, therefore, spoke of the phenomenon as "anaphylaxis" ("action anaphylactique de certains venins") to express its antithesis to prophylaxis or protective effects.

Although it has been disputed by a number of writers that Richet's investigations constitute the beginnings of our modern understanding of the anaphylactic phenomena, yet his recognition of the distinct dependence of the hypersusceptible condition upon a preceding inoculation with the same substance, and his conclusion that a definite incubation time must elapse after the first injection before susceptibility is developed, defined two of the most important criteria of the condition and initiated purposeful investigations in this field. It is true, on the other hand, that, like v. Behring and most of his other predecessors, he was working with primarily toxic substances, and the final recognition of the general biological significance of the anaphylactic phenomenon was necessarily deferred until a similar development of hypersusceptibility was noted in animals injected with various antigens which of themselves were entirely harmless. In this the history of anaphylactic investigations is similar to that of other reactions to antigen injections, lysin, agglutinin, and precipitin formation, in which the first observations were made upon pathogenic bacteria or their products, and in which subsequent extension of the investigations revealed that the response to inoculation with bacterial proteins represented merely a single phase of a general biological reaction on the part of animals to treatment with the large class of substances known as antigens.

³ Richet and Portier. *C. R. de la Soc. Biol.*, p. 170, 1902.

⁴ Richet and Héricourt. *C. R. de la Soc. Biol.*, 1898.

⁵ Portier and Richet. *C. R. de la Soc. Biol.*, p. 170, 1902.

This generalization of Richet's observations had really been foreshadowed by the observations of Magendie and by the experiments of Flexner quoted above, but this work had been lost sight of and the attention of investigators was again focused upon the problem mainly by the publication of Arthus⁶ in 1903 on the repeated injection of horse serum into rabbits, and some observations made upon guinea pigs by Theobald Smith and communicated by him in 1904 to Ehrlich.

Arthus⁷ found that horse serum injected into rabbits by any of the usual paths of entrance is entirely innocuous. It is possible to inject 10, 20, or even 40 c. c. without harm. If, however, one repeatedly injects small amounts, 5 c. c. or less, subcutaneously, at intervals of several days, eventually the later injections will give rise to infiltrations, edema, sterile abscesses, and even gangrene at the points of injection. He recognized that this was not due to cumulative action, and that it was not necessary to inject several times in the same place to produce the characteristic response. For instance, the early injections might be made into the peritoneum, the subsequent ones into the skin, and the local reactions to the later injections might nevertheless ensue. In other words, he recognized the systemic nature of the phenomenon and regarded it as analogous to the observations of Richet in that he spoke of the hypersensitive rabbits as "anaphylactisés" by a series of preparatory injections.

The "phenomenon of Theobald Smith" is closely related to that of Arthus, and was made in the course of the standardization of diphtheria antitoxin in guinea pigs. It was noticed that guinea pigs which had been used for this purpose and had survived had acquired great susceptibility to subsequent injections of normal horse serum made several days or weeks later.

With these observations as points of departure, together with the studies of v. Pirquet and Schick⁸ upon the clinical manifestations of antitoxin injections into human beings, a number of investigators took up the problem, chief among them Rosenau and Anderson, of the United States Hygienic Laboratory, and R. Otto, of the Frankfurt Institute of Experimental Therapy.

Although the paper of Otto⁹ appeared in print a little earlier than did the first one of the American workers, the investigations were independent and almost synchronous. Their results, moreover, confirm each other in all essentials. Otto showed that the Theobald Smith phenomenon was entirely independent of the toxin or anti-

⁶ Arthus. *C. R. de la Soc. Biol.*, Vol. 55, p. 817, Réunion biol., Marseille, June, 1903.

⁷ Arthus et Breton. *C. R. de la Soc. Biol.*, 55, p. 1478.

⁸ Von Pirquet u. Schick. "Die Serumkrankheit," Deuticke, Wien, 1906.

⁹ Otto. "Das Theobald Smithsche Phaenomen, etc., v. Leuthold Gedenkschrift," Vol. 1, 1905; also Otto in *Ergänzungsband 2*, "Kolle u. Wassermann Handbuch," etc.

toxin contents of the injected serum, but could be produced (though somewhat less markedly) with horse serum alone. He also showed that, while a preliminary injection of horse serum "sensitized" a guinea pig to a subsequent dose given after an interval of 10 to 12 days, the repeated injection of considerable quantities at short intervals produced a condition of "antianaphylaxis" or immunity to the later injections. Otto, too, excluded from his results the direct relation of the anaphylactic state with the possible presence of serum precipitins, a thought suggested by Morgenroth in his interpretation of the observations of Magendie mentioned above.

Rosenau and Anderson¹⁰ had attacked the problem with the primary purpose of throwing light upon the occasional accident of sudden death following the injection of diphtheria antitoxin into human beings. Since the detailed description of their extensive investigations would tend to render more difficult the exposition of an already sufficiently complicated subject, it will be best to tabulate the chief results of this classical series of their earlier papers. Briefly, these are as follows:

1. A single injection of horse serum into guinea pigs, harmless in itself, renders these animals hypersusceptible to a subsequent injection given after a definite interval or incubation time.

2. This interval, with the ordinary dosages employed (about 1 to 2 c. c.), was about 10 days. Properly carried out injections after this period were usually fatal.

3. The known antibodies, antitoxins, hemolysins, and precipitins, are not responsible for the reaction.

4. The reaction is "quantitatively" specific, injections of horse serum sensitizing to horse serum only. (The question of specificity will be further discussed below.)

5. The sensitive condition is transmissible from mother to offspring,¹¹ the young of sensitized mothers being hypersusceptible to a *first* injection of horse serum.

6. The reaction is extremely delicate. Rosenau and Anderson succeeded in sensitizing in one case with 0.000001 c. c. (one one-millionth) of horse serum.

7. The hypersusceptible state is not a transient condition, but may last a long time.

8. Sensitization, or the production of the hypersusceptible condition, can be carried out, not only with the various animal and vegetable proteins employed in the first experiment, but can be brought

¹⁰ Rosenau and Anderson. *U. S. Pub. Health and M. H. S. Hyg. Lab. Bull.* 29, 1906; 30, 1906; 36, 1907; *Journ. Med. Res.*, Vol. 15, 1906, Vol. 16, 1907; also *Jour. Inf. Dis.*, Vol. 4, 1907, Vol. 5, 1908.

¹¹ It is important practically, as Anderson points out, that a female guinea pig may transmit to its young sensitiveness to horse serum and immunity to diphtheria toxin.

about by the use of extracts of various bacteria. In such cases also the reaction is specific. The first determinations with bacterial extracts carried out by Rosenau and Anderson were made with colon, anthrax, typhoid, and tubercle bacilli.

By these observations, then, the possibility of a direct relation between the phenomena of anaphylaxis and infectious diseases in animals was indicated.

This, in essence, is the harvest of the two earliest purposeful researches into this problem. A large number of investigators now took up the question, and its further elucidation, as we shall see, has proved, not only the most directly fruitful of the phases of recent immunological studies, but has thrown much indirect light upon antigen-antibody reactions apart from the anaphylactic phenomena themselves.

Before entering into the further discussion of the experimental data, however, it will be necessary to describe briefly the clinical manifestations which follow upon the second injection of an anaphylactic antigen into a sensitized animal, manifestations which we have heretofore summarized in the phrase "anaphylactic shock." For there has been much controversy regarding the physiological mechanism which lies at the bottom of these symptoms, and the matter has been complicated by the unquestionably different reactions occurring in various species of animals in response to the anaphylactic experiment.

Since anaphylactic studies were begun largely as the result of Theobald Smith's observations upon guinea pigs, and subsequent study has revealed these animals as peculiarly susceptible to the anaphylactic poison, the large bulk of the experimental data at our disposal was worked out upon these animals. In consequence our understanding of the mechanism of the reaction is based largely upon guinea pig studies.

If a properly sensitized guinea pig receives a second injection of an antigen after a suitable incubation time a very characteristic train of symptoms ensues. There is usually a short preliminary period—lasting either a fraction of a minute or several minutes according to the violence of the reaction and the mode of administration—during which the pig appears normal. At the end of this time the animal will grow restless and uneasy, and will usually rub its nose with its forepaws. It may sneeze and occasionally emit short coughing sounds. At the same time an increased rapidity of respiration is noticeable and the fur will appear ruffled. In light cases the animals may remain in this condition, with further irregularity and difficulty of respiration, possible discharges of urine and feces; then gradual slow recovery may set in, with complete return to normal in from 30 minutes to several hours. In more severe cases these preliminary stages are rapidly followed by great apparent

weakness. The animals fall to the side, the legs and trunk muscles twitch irregularly, and the respiration becomes slow and shallow; the thorax never entirely contracts, but remains in a more or less expanded condition. The very evident dyspnea is of an inspiratory character. The excursions of the lung itself seem to grow shallower and shallower in spite of apparent strong inspiratory efforts—the volume of the thorax and lung remaining in the expanded condition. At this stage evidences of motor irritation may appear, in that the animal may arise and attempt to run. More often, however, in this phase general convulsions set in, often several times repeated, and in these the animals usually die.

On the other hand, after cessation of convulsions they may lie perfectly still on the side as though paralyzed, the breathing becoming gradually slower and more shallow, finally ceasing entirely. The heart may continue to beat for a considerable time after the breathing has stopped.

If such an animal is immediately autopsied a very characteristic condition is found—to which, in the essentials, attention was first called by Gay and Southard.¹² They speak of finding “pulmonary emphysema as a constant feature at autopsy,” and attribute the anaphylactic death in guinea pigs to cessation of respiration in the inspiratory phase under the influence of respiratory central intoxication.

The lungs of such guinea pigs after death are found distended and completely filling the thorax. They are usually pale and bloodless and do not collapse as the pleuræ are opened. On microscopic examination the alveoli are seen to be distended and small hemorrhages may appear upon the serous surfaces. According to Gay and Southard, furthermore, histological study of the other organs shows also hemorrhages in the brain, stomach, heart, cecum, and spleen—more rarely in other organs, and there are local fatty changes in the capillary endothelium which they regard as causatively related to the hemorrhages.

That the respiratory symptoms are the most striking feature of the clinical picture of guinea pig anaphylaxis had, as a matter of fact, been noticed by Rosenau and Anderson. A detailed physiological study of the mechanism of the respiratory death in these cases was first made, however, by Auer and Lewis¹³ in 1909.

These investigators showed that, during the later respiratory symptoms, little or no air enters the lungs, although the animal makes violent respiratory efforts. This is due, as they found, to a tetanic contraction of the small bronchioles, which practically oc-

¹² Gay and Southard. *Jour. Med. Res.*, Vol. 16, 1907; Vols. 18 and 19, 1908.

¹³ Auer and Lewis. *Jour. of the A. M. A.*, Vol. 53, p. 458, 1909; *Jour. Exp. Med.*, Vol. 12, 1910.

cludes the air passages. That the origin of this contraction is not, as previously supposed, of central origin, but is referable to peripheral cause, they proved by showing that the same phenomena occur in the guinea pigs even after the cord and medulla have been destroyed and the vagi divided. In such cases, of course, with the cord and medulla destroyed, artificial respiration had to be done, and when the symptoms set in it was found that the lungs could no longer be expanded by the same force of artificial respiration which before this had been sufficient.

They showed also that the non-collapsible expansion of the lungs after death was due to imprisonment of the air in the alveoli by the contracted musculature of the small bronchioles, and further confirmed their opinion of the peripheral origin of this contraction by the important discovery that atropin will markedly protect, often preventing death or hastening recovery. It is noteworthy, too, that Auer and Lewis speak of occasionally finding slight pulmonary edema, a feature which Biedl and Kraus consider incompatible with true anaphylaxis.

Anderson and Schultz,¹⁴ who have confirmed much of the work of Auer and Lewis, find that not only atropin will prevent asphyxiation in these cases, but methane, chloral hydrate, adrenalin, and pure oxygen will exert a similar effect. The animals may be saved from suffocation in this way, but may nevertheless die, probably as the result of lowered blood pressure.

The observations of Auer and Lewis have been further confirmed especially by Biedl and Kraus,¹⁵ who regard it as well established that anaphylactic death in guinea pigs is caused primarily by suffocation, due to tetanic spasms of the musculature of the small bronchi. These spasms are not of central origin, but are peripherally initiated, possibly by direct action upon the smooth muscle itself. The fact that atropin is not effective in preventing death in all severe cases is no argument against this, since such an effect would naturally depend upon the relation between the amount of atropin given and the severity of the attack. In this connection the studies that have been made upon the irritability of smooth muscle fibers in normal and in sensitized animals are of great interest. Schultz,¹⁶ following out an observation made by Rosenau and Anderson, studied the intestinal muscle of normal sensitized guinea pigs excised and suspended in Howell's solution. In this way he showed that during the period of hypersusceptibility the smooth muscle is abnormally sensitive to treatment with the antigen. The contraction which normally occurs

¹⁴ Anderson and Schultz. *Proc. Soc. Exp. Biol. and Med.*, 7, 1909, p. 32.

¹⁵ Biedl und Kraus. *Zeitschr. f. Immunitätsforschung*, Vol. 7, 1910; *Centralbl. f. Physiol.*, 1910; *Wien. klin. Woch.*, No. 11, 1910.

¹⁶ Schultz. *Jour. Pharm. and Exp. Therap.*, 1, 1910; 2, 1910.

in smooth muscles under the influence of serum is markedly augmented if the preparations are taken from sensitized animals.

In addition to these predominant features of the anaphylactic symptomatology in guinea pigs, there are a number of secondary reactions which, though less prominent, are nevertheless of considerable interest and theoretical importance. The conditions in the circulation are probably, to a great extent, dependent upon the respiratory condition, and the fall of blood pressure in guinea pigs is regarded by some investigators as merely a secondary manifestation just preceding death. The fall of temperature first described by H. Pfeiffer,¹⁷ however, seems to be an occurrence which, though standing in no causative relation to the symptoms as a whole, is so constant and well marked that it has been taken by a number of workers as one of the necessary criteria for the characterization of the anaphylactic condition.

There is, indeed, an almost regular drop of several degrees in the rectal temperature, and a close observation of this may be of much aid in determining the occurrence of mild reactions, when other symptoms of shock are not strongly marked. Pfeiffer¹⁸ himself goes so far as to claim that by this symptom alone delicate anaphylactic reactions may be determined when all other symptoms are lacking.

Friedberger,^{19 20} too, has found the sudden drop of temperature a very regular occurrence, and has employed this method of study for the analysis of the intensity of anaphylactic shock. He calls attention to the apparent difference between infection and anaphylaxis in this respect in that in the former there is fever, in the latter there is depression of body heat; but, at the same time, he points out that this discrepancy is an apparent one only, and determined by quantitative differences, for when he treated sensitized animals with varying doses of antigen he found that quantities which produced other anaphylactic symptoms of noticeable degree would regularly depress the temperature as Pfeiffer had shown. It was possible, however, to determine a minimal dose necessary for temperature reduction. Quantities just below this left the temperature unchanged, and still smaller quantities produced fever or even increased the temperature. This fact is extremely significant in that, as we shall see, it has an important bearing upon views which interpret bacterial infection as a series of anaphylactic poisonings, the multiplying bacteria furnishing the constant supply of minute amounts of antigen. This thought, indeed, based also on the study of

¹⁷ H. Pfeiffer. *Wien. klin. Woch.*, No. 1, 1909.

¹⁸ Pfeiffer u. Mita. *Zeitschr. f. Immunitätsforschung*, Vol. 4, 1910.

¹⁹ Friedberger. *Deutsche med. Woch.*, No. 11, 1911.

²⁰ Friedberger und Mita. *Zeitschr. f. Immunitätsforschung*, Vol. 10, 1911.

temperature curves in animals, was expressed by Vaughan²¹ as early as 1909, and was developed by him with Cumming and Wright²² in an extensive study upon what he called "protein fever." It was shown in these experiments that continued fever, not unlike that of infectious diseases, could be produced in rabbits by repeated subcutaneous injections of primarily harmless substances, such as egg white and vegetable proteins. The conditions observed and the conclusions drawn from them in this work, as well as in the similar investigations of other workers, were clearly foreseen by Vaughan in his early investigations on proteid split-products studies, which we will find occasion to discuss in a later section.

The rigidity of the diagnostic value of the temperature relations for anaphylactic shock in particular, as advanced by Pfeiffer, was somewhat weakened by Ranzi's²³ observations that foreign serum may produce temperature depression when injected into perfectly normal animals and that, injected into sensitized animals, the same reaction may follow if other proteins than the original antigen were administered.

Although these objections of Ranzi are perfectly just, yet there is such a marked quantitative difference between the reaction in normal and in sensitized animals that, in principle, Pfeiffer's claim is not invalidated. Friedberger²⁴ very logically remarks that, after all, the phenomena of sensitization as well as those of immunity are merely an exaggeration of normal physiological conditions, and in experiment he has shown that, whereas noticeable depressions of temperature will follow in the normal animal only upon quantities of antigen exceeding 0.5 c. c., the temperature of the sensitized animal may be depressed by amounts as small as 0.0005 c. c.

Apart from the symptoms so far discussed, there are other less apparent characteristics of anaphylaxis in guinea pigs, all of which, however, possess considerable importance theoretically. The most significant of these is the reduction in the amount of alexin or complement, first noticed by Sleeswijk,²⁵ which occurs after the injection of the second or toxogenic dose—during the development of shock. This phenomenon is so closely interwoven with the later theoretical aspects of anaphylaxis that we will defer its discussion until we have completed a more general survey of the field.

In guinea pigs, as in dogs, Friedberger and others have also seen a lowered coagulability of the blood and a temporary diminution of the polynuclear leukocytes (leukopenia) during shock.

²¹ Vaughan. *Zeitschr. f. Immunitätsforschung*, Vol. 1, 1909.

²² Vaughan, Cumming, and Wright. *Zeitschr. f. Immunitätsforschung*, Vol. 9, 1911.

²³ Ranzi. *Zeitschr. f. Immunitätsforschung*, Vol. 2, 1909; *Wien. klin. Woch.*, No. 40, 1909.

²⁴ Friedberger u. Mita. *Loc. cit.*

²⁵ Sleeswijk. *Zeitschr. f. Immunitätsforschung*, Vol. 2, 1909.

During the earlier periods of experimentation there was a marked discrepancy in the ease with which guinea pigs could be sensitized by American and German investigators, on the one hand, and by Besredka and Steinhart in France, on the other. The mortality, upon second injection, was much higher, with like quantities of horse serum in the hands of the first-named. In attempting to explain this, Rosenau and Anderson carried out typical experiments with horse serum sent to them by Besredka and obtained high percentages of fatal results. They believe, for this reason, that the differences cannot be accounted for by variations in the toxicity of the horse sera, but conclude that probably there are varying grades of susceptibility to the reaction in guinea pigs of different breeds.

Next to guinea pigs the animals most commonly employed for anaphylactic experiment are rabbits and dogs. In both of these the symptoms and autopsy findings differ markedly from each other and from those observed in guinea pigs.

In sensitized rabbits the injection of a second dose of the antigen is usually followed, after a short but definite incubation time, by great weakness with, often, discharge of urine and feces. The animals sink down until the abdomen touches the ground, the legs are stretched out weakly but not paralyzed, and the head may drop forward or to one side. After this, the animal may gradually fall upon its side and lie motionless except for labored and irregular breathing and occasional twitching of the legs and head. Sometimes this gradual relaxation may be interrupted by a sudden motor irritation, the rabbit suddenly getting up and running a short distance but soon falling down again apparently from a sudden return of the muscular weakness. During these running spells it seems as though there was no sense of direction or purpose—the animals running into obstructions or off tables as the case may be. During this period general convulsions and a drawing back of the head by a tetanic spasm of the muscles of the neck are not uncommon. Death may occur within a few minutes, or it may follow a gradually increasing weakness in the course of several hours. The fall of blood pressure here seems to be purely secondary to the general failure of all the functions.²⁶

Anaphylaxis in dogs has been very extensively studied, especially by Biedl and Kraus,²⁷ and by Pearce and Eisenbrey.²⁸ The symptoms in dogs are characterized by a rapid progressive fall in the blood pressure, followed by the symptoms of cerebral anemia. Anaphylactic dogs, after injection, will at first grow restless, vomit, and

²⁶ Arthus. *Arch. Internat. de Physiol.*, 7, 1909.

²⁷ Biedl and Kraus. *Loc. cit.*; also in "Kraus u. Levaditi Handbuch," *Ergänzungsband* 1.

²⁸ Pearce and Eisenbrey. *Proc. Soc. Exp. Biol. and Med.*, 7, 1909, p. 30; *Transact. Congr. Am. Ph. and S.*, Vol. 8, 1910.

pass urine and feces. They then grow rapidly weak, fall to the ground, and continue to twitch and vomit and the respiration becomes labored and irregular. There is general weakness of the muscles, but no paralysis. The marked, constant, and characteristic feature of the condition in these animals is the fall of blood pressure. There is also a lessened coagulability of the blood, much more strongly developed than in guinea pigs and rabbits.

According to Biedl and Kraus this may amount to almost a prevention of the coagulation in anaphylactic dogs.

As in other animals the blood picture is changed in that there is a falling off of the total number of leukocytes with a relative diminution of polynuclear cells.

Quantitative measurements by Calvary,²⁹ moreover, have shown that anaphylaxis in dogs is accompanied by a marked increase of the lymph flow (7 times the amount observed in normal dogs in the same time) and, by controlling the blood pressure with barium chlorid, that this lymphagogue action is not directly dependent upon the low pressure. This observation is of especial interest in connection with the similarity of anaphylaxis to peptone poisoning in which Heidenheim³⁰ noticed a similar increase of the lymph.

Pearce and Eisenbrey found, at autopsy of dogs dead of anaphylactic shock, subserous petechial hemorrhages in the rectum and gall bladder, hemorrhagic spots on the gastric and duodenal mucosa, and in the colon. According to these workers, in agreement with Biedl and Kraus, the fall of blood pressure is not due to central causes but depends upon influences exerted upon the peripheral vasomotor system. Biedl and Kraus believe that this action is exerted upon the muscle cells themselves rather than on the nerve endings. They admit the inconclusiveness of their experimental data, but take the above standpoint because of the fact that adrenalin, which acts by stimulation of the vasomotor nerve endings particularly, does not raise the low pressure in dogs during anaphylaxis while barium chlorid, which acts upon the smooth muscle fibers themselves, strongly raises the blood pressure in such animals. Pearce and Eisenbrey are inclined to believe that the action is chiefly upon the nerve endings, though both factors, nerve and muscle, may be involved. They worked with apocodein, a substance which, in large doses, paralyzes the vasomotor nerve terminals.³¹

When a sensitized dog was treated with apocodein and the antigen then injected, no further drop of pressure was obtained. Apparently a paralysis of the vasomotor nerve endings had removed the point of attack upon which the anaphylactic poison could act.

In addition to the symptoms already enumerated Weichhardt and

²⁹ Calvary. *Münch. med. Woch.*, No. 13, 1911.

³⁰ Heidenheim. *Pflüger's Archiv*, 49, 1891.

³¹ Brodie and Dixon. *Jour. of Phys.*, 30, 1904.

Schittenhelm³² claim that anaphylaxis in dogs is invariably accompanied by a severe local reaction in the gut. The intestinal mucosa is swollen and contains miliary hemorrhages and the lumen is often filled with a mucus mixed with blood. In the further analysis of the anaphylactic reaction in dogs, Manwaring³³ has recently reported observations of great interest. He investigated the participation in anaphylactic shock of the various organs and determined that shock did not occur when the abdominal vessels were ligated just above the diaphragm. In further localizing the source of shock he found that exclusion of the spleen, stomach, kidneys, suprarenals, and ovaries from the circulation had no effect upon the occurrence of anaphylactic shock. However, when he operated in such a way that the liver was thrown out of circulation, none of the seven dogs that he used reacted with anaphylactic shock to the injection of serum. He concludes from this that the liver is directly responsible in some way for the production of anaphylaxis. The intestines, too, were found, by a similar procedure, to take part, though to a less important extent than the liver.

Other animals than those mentioned have been little used for anaphylactic experiment. Observations incidental to other work, however, have shown that horses and goats are particularly sensitive. In goats the writer has observed both serum and bacterial anaphylaxis, and the symptoms here were those of general trembling, weakness, labored respiration, and involuntary evacuation of urine.

The occurrence of anaphylaxis in man will be discussed in a subsequent section.

The manifestations of "active anaphylaxis," therefore, consist in the profound physiological changes occurring in animals when re-injected after a definite interval with certain substances which, on first injection, were practically harmless. The factors which are of fundamental importance in determining the development of this hypersusceptible or anaphylactic state consist in the nature of the injected substance, the quantity injected, and the interval between administrations. To a great extent, too, the violence of the reaction is dependent upon the path by which the particular substance enters the body.

Each of these factors, therefore, requires detailed consideration before we can intelligently proceed with a further analysis of the condition.

The substances with which animals may be sensitized are, in all particulars, identical with the class of substances which we have characterized as "antigens." In fact, up to the present time, there has not been a single authenticated exception to this, and from our present understanding of the mechanism of anaphylaxis we may

³² Weichhardt and Schittenhelm. *Deutsche med. Woch.*, 19, 1911.

³³ Manwaring. *Zeitschrift f. Immun.*, Vol. 18, 1911.

safely predict that no such exceptions will be found. It is the large class of proteins, therefore, whatever their source, which may act as the "anaphylactic antigens." However, in this connection as well as in the larger problem of the nature of antigens in general, it has been difficult to decide whether or not the antigenic property is entirely confined to proteins or whether other substances, such as the lipoids, must be included in the definition. The problem has been the same here as in other serum phenomena, but much special experimentation has been done upon the question with particular reference to anaphylaxis and the possibility of sensitizing animals with lipoids.

As in the case of similar investigations in regard to antibody formation, the results obtained in this work have been somewhat confusing. Pick and Yamanouchi³⁴ extracted beef and horse sera with alcohol, and evaporated and redissolved the solutions until they neither contained coagulable protein nor gave the Biuret reaction. With this material they obtained a few positive anaphylactic experiments. Similarly curious are the results of Bogomolez,³⁵ who succeeded in sensitizing and producing shock with the lipoids extracted from egg yolks. Although such experiments would tend to persuade us that lipoidal substances may actually have sensitizing (therefore antigenic) functions, this does not follow necessarily. As Pick and Yamanouchi themselves point out, it is practically impossible to demonstrate with certainty the presence of slight traces of proteins as impurities in lipoid preparations, and we know especially from Rosenau and Anderson's work how minute are the quantities of antigen which still serve to sensitize. It is possible, moreover (a thought developed particularly by Pick and Schwartz³⁶ and by Landsteiner³⁷), that we are dealing in many cases with combinations of protein and lipid—a form of chemical substances of which very little is known analytically, but the existence of which many biological facts lead us to assume.

That the anaphylactic reaction is specific we have mentioned in the brief summary we have given of Rosenau and Anderson's work. These authors use the adjective "quantitative," by which they simply mean to convey that the specificity here is not absolute, any more than it is absolute in the case of any of the known serum reactions. An animal sensitized with a certain variety of protein, animal serum, etc., reacts with disproportionately greater delicacy to a second injection of the same variety than of any other substance. In fact, apart from a few cases mentioned by Gay and Southard, there are not many instances of marked non-specific anaphylactic reactions. Still we

³⁴ Pick and Yamanouchi. *Zeitschr. f. Immunitätsforschung*, Vol. 1, 1909.

³⁵ Bogomolez. *Zeitschr. f. Immunitätsforschung*, Vols. 5 and 6, 1910.

³⁶ Pick and Schwartz. *Biochem. Zeitsch.*, 15, 1909.

³⁷ Landsteiner. Ref. "Weichhardt's Jahresbericht," 6, 1910.

would expect here, as in other serum reactions, a certain limitation in the degree of specificity, and Otto recommends the less delicate subcutaneous method of testing for all experiments in which questions of specificity are involved. This point we will touch upon a little later.

An interesting addition to our knowledge of such specificity was made by experiments of Rosenau and Anderson,³⁸ which showed that a guinea pig could be rendered sensitive at one and the same time to blood serum, eggwhite, and milk, reacting specifically to each on second injection.

In anaphylaxis, again analogous to antibody reactions in general, the specificity, as a rule, is one of species. In other words, the protein of any animal is specific for the proteins of its particular species generally, there being definitely similar characteristics in the body proteins of animals of like species which, though chemically indefinable, are nevertheless delicately determinable by biologic reactions. In considering specificity of precipitins, however, we have seen that there are exceptions to the specificity of species expressed in the phenomenon of so-called organ specificity. The same thing has been shown for anaphylaxis. Kraus, Doerr, and Sohma³⁹ were able to show that animals sensitized with protein from the crystalline lens were hypersusceptible to lens protein generally, whether this came from the species from which the original lens was taken, or whether some other variety of animal had furnished it. On the other hand, animals so sensitized, while hypersusceptible to lens protein generally, did not react to injections of homologous blood.⁴⁰ In other words, this organ contains a characteristic variety of antigen (protein) peculiar to this kind of organ throughout the different animal species, but not common to other tissues and organs of the same animal. Results similar to these were obtained by von Dungern and Hirschfeld⁴¹ in the case of testicular protein, although here the phenomenon seemed to be less rigidly organ-specific than in the preceding case. These writers worked not with the systemic anaphylactic reaction, but with the localized (allergie) reaction, described above as the phenomenon of Arthus. They injected extracts of the testicular materials into the ears of rabbits and incidentally made the very curious observation that pregnant females would not infrequently react to a first injection without previous sensitization.

Of great importance also in connection with the subject of organ

³⁸ Rosenau and Anderson. *Jour. Inf. Dis.*, Vol. 4, 1907.

³⁹ Kraus, Doerr, and Sohma. *Wien. klin. Woch.*, No. 30, 1908.

⁴⁰ Andrejew. *Arb. a. d. kais. Gesundh. Amt.*, Vol. 30, 1909.

⁴¹ Von Dungern and Hirschfeld. *Zeitschr. f. Immunitätsforschung*, 4, 1910.

specificity is the further discovery by Uhlenhuth and Haendel⁴² that animals can be sensitized with their own lens protein, a fact which opens the possibility of other forms of "autosensitization" and consequently of much opportunity for clinical speculation. Rosenau and Anderson,⁴³ indeed, have found that guinea pigs can be sensitized by means of extracts of guinea pig placenta. They have applied this to the possible explanation of eclampsia, and similar reasoning, as we shall see, has been utilized in many other conditions. Attempts have also been made to show, by the anaphylactic reaction, that the tissue of malignant tumors possesses such "tissue-specific" or "organ-specific" qualities. Yamanouchi,⁴⁴ indeed, claims to have shown this, but his results were not confirmed by Apolant,⁴⁵ and the writer has carried out a series of entirely negative experiments upon the same subject. However, in view of the great difficulty of obtaining any kind of anaphylactic reaction in mice, the animals in which the tumor experiments were carried out, there is little information to be obtained from negative results of this kind.

The delicate quantitative method of studying problems of specificity, which the reaction of anaphylaxis supplies, has further served to revive the unsettled question of the "organ-specific" properties of the tissues of such organs as the liver, spleen, kidney, blood, etc. Indeed, Pfeiffer⁴⁶ has published results which would seem to encourage the belief of the existence of such specificity. However, Ranzi⁴⁷ had previously obtained entirely negative results, and Pearce, Karsner, and Eisenbrey⁴⁸ have recently made a careful inquiry into the same problem with similar failure to determine such organ-specificity.

In this, then, as well as in other respects, the substances by which animals may be sensitized are entirely similar to antigens in general.

The substances which sensitize, therefore, are those which have the property of antibody formation, a statement self-evident from what has been said before, but which is again emphasized because of its very important bearing upon later theoretical considerations.

As in antibody production, variations in experimental anaphylaxis are, to some extent, dependent upon the manner in which the antigen is introduced into the body. It is now well known that sensitization may be accomplished by a first injection given subcutaneously, intravenously, intraperitoneally, or intrapleurally. At the second or toxogenic administration shock may probably be best

⁴² Uhlenhuth and Haendel. *Zeitschr. f. Immunitätsforschung*, 4, 1910.

⁴³ Rosenau and Anderson. *U. S. Pub. Health and M. H. S. Hyg. Lab. Bull.* 45.

⁴⁴ Yamanouchi. *C. R. de la Soc. Biol.*, Vol. 66, 1909, p. 754.

⁴⁵ Apolant. *Zeitschr. f. Immunitätsforschung*, Vol. 3, 1909.

⁴⁶ Pfeiffer. *Zeitschr. f. Immunitätsforschung*, Vol. 8, 1910.

⁴⁷ Ranzi. *Zeitschr. f. Immunitätsforschung*, Vol. 2, 1909.

⁴⁸ Pearce, Karsner, and Eisenbrey. *Jour. Exp. Med.*, Vol. 14, 1911.

induced and with the smallest quantities by the intravenous method. Besredka and Steinhardt,^{49 50 51 52} who began their studies soon after the first publications of Rosenau and Anderson, came to the conclusion that the most effectual and rapid method of producing the anaphylactic shock consisted in direct injection into the brain. Curiously enough, while Besredka and Steinhardt obtained the most violent reactions by injection of the second or toxogenic dose into the brain, they were unable to sensitize by this path, at least with doses of 1/4000 c. c., which sufficed to sensitize by the intravenous method. Rosenau and Anderson, in repeating this work, obtained similar results with very minute amounts, but found that intracerebral sensitization could be accomplished by doses of 0.0001 c. c., or more. According to them, animals intracerebrally sensitized became anaphylactic more rapidly than those in which the injections were subcutaneous. In the former the incubation time was about 7 days, while in the latter it was never less than 9. Lewis,⁵³ in his thorough study on the same subject, made extensive use of the direct intracardial method of injection. In other words, any method of introducing the foreign protein into the blood or tissues seems to lead both to sensitization and to toxic effect, and those methods which introduce the substance, on reinjection, directly into the blood stream or the brain induce the most violent symptoms with the relatively smallest dosage. According to Otto and others, the subcutaneous method, while followed by less violent symptoms, is the method to be preferred when questions of specificity are involved, for, while the reaction is specific in the ordinary sense, yet it is extremely delicate and therefore, as Rosenau and Anderson put it, "quantitatively specific." The less violent subcutaneous method, therefore, might be said to have the same purpose here that dilution of the antigen or immune serum has in safeguarding against error when carrying out specific precipitin or agglutinin reactions.

Whether or not sensitization can be accomplished by introduction of the antigen into the intestinal canal, feeding, in other words, is still to some extent an open question and of great importance in view of the many clinical manifestations (urticaria, albuminuria, etc.) which are attributed to possible individual hypersusceptibility to certain proteins taken in the diet (idiosyncrasies). Rosenau and Anderson, in their earliest paper, report success in sensitizing guinea pigs by the feeding of horse meat and horse serum. McClintock and

⁴⁹ Besredka and Steinhardt. *Ann. de l'Inst. Past.*, p. 117, 1907; *ibid.*, p. 384.

⁵⁰ Besredka. *Ibid.*, pp. 777, 950, 1907; *ibid.*, p. 496, 1908; p. 166, 1909; p. 801, 1909.

⁵¹ Also: *Bull. de l'Inst. Past.*, Nos. 19, 20, 21, 1908; No. 17, 1909.

⁵² Also: *C. R. de la Soc. Biol.*, p. 478, 1908, Vol. 65; p. 266, 1909, Vol. 67.

⁵³ Lewis. *Jour. Exp. Med.*, Vol. 10, 1908.

King⁵⁴ failed to confirm this, and the observations of other writers seem to bear them out. However, when we consider that Ascoli, Oppenheimer, and others have shown that proteins fed to animals in large quantities may be subsequently demonstrated not only in the circulating blood but occasionally even in the urine by means of the precipitin reaction, there seems to be little room for doubting that antigen may enter the circulation unchanged, though possibly only under abnormal local conditions of the intestine. This, together with Rosenau and Anderson's demonstration of the extremely small amount of antigen necessary to sensitize, furnishes all the conditions necessary for anaphylaxis by way of the intestinal canal.

A study made by Lesné and Dreyfus⁵⁵ seems to us to have explained the contradictory results of other workers on this phase of the problem. Without being able to associate the destruction of the sensitizing function with either the gastric or pancreatic secretions, they were nevertheless successful in showing that sensitization could be carried out regularly if the antigen were injected after laparotomy into the large intestine, whereas similar injections into the stomach or small intestine were negative. In these experiments we must take into consideration that the conditions following laparotomy, such as temporary intestinal atony and congestion, may have exerted considerable influence upon the positive outcome of their large intestine injections. Whereas they do not, therefore, permit us to assume the possibility of sensitization through the normal alimentary canal, they nevertheless confirm the assumption of the possibility of sensitization by this path under the influence of slightly abnormal local conditions.

In this connection Besredka's⁵⁶ experiments on the production of anti-anaphylaxis by the intestinal administration of protein are of interest. He found that, if sensitized animals were given 5 c. c. of the antigen (milk) by rectum, they were thereby protected from the reaction following in controls upon a second injection. In his later experiments with egg white it appeared that the protection could also be conferred by mouth, but that in this case it developed more slowly, it being necessary to wait two days after ingestion before the anti-anaphylaxis had developed sufficiently to protect. Since attempts by mouth were not as rapidly successful as those *per rectum*, it is clear that these facts are in keeping with Lesné and Dreyfus' results in showing that the antigen is probably absorbed chiefly or solely from the large intestine. Lesné and Dreyfus sensitized by way of the intestine, and administered the second or toxogenic dose intravenously, and since, as we shall see, minute doses may

⁵⁴ McClintock and King. *Jour. Inf. Dis.*, 3, 1906. See section on normal antibodies.

⁵⁵ Lesné and Dreyfus. *C. R. de la Soc. Biol.*, Vol. 70, p. 136, 1911.

⁵⁶ Besredka. *C. R. de la Soc. Biol.*, Vol. 65, 1908; Vol. 70, 1911.

suffice to sensitize, whereas 100 or more times this amount is necessary to produce intoxication, it is easy to understand why the rectal route sensitized in Lesné and Dreyfus' work, but no toxic effects followed in the experiments of Besredka. Furthermore, the slow absorption from the intestine in these experiments explains the development of anti-anaphylaxis in Besredka's work, in that they are, in this respect, analogous to later experiments of Friedberger, cited below, in which it was shown that sensitized guinea pigs, which could (in controls) be killed by rapid intravenous injection of 0.1 c. c. of antigen and less, would withstand many times this amount when gradually administered by slow injections extending over periods of an hour or longer.

In referring to the quantities of antigen by which sensitization may be accomplished, we have already called attention to the very small amounts which have been found sufficient for this purpose. There seems, indeed, to be a wide latitude in this regard, the required quantities ranging from as little as a millionth of a cubic centimeter (Rosenau and Anderson) to as much as 10 c. c. or more. On second injection, however, toxic effects are never produced by quantities as minute as those which suffice for sensitization, though here, too, a wide range of effectual amounts exists. An important problem, moreover, is the relation which has been said to exist between the sensitizing dose and the interval necessary for the development of the hypersusceptible state (anaphylactic incubation time). In their first publications, Rosenau and Anderson, Otto, and others expressed the opinion that the length of incubation time was inversely proportionate to the size of the sensitizing dose; in other words, animals sensitized with small quantities (0.01 c. c. or less) would become hypersusceptible and react to a second injection in from 8 to 12 days, whereas animals receiving two, three, or more cubic centimeters of the antigen would take weeks or months to become anaphylactic. The same opinion was expressed by Otto,⁵⁷ and is now generally found in the literature. Later experiments of Rosenau and Anderson,⁵⁸ however, have seemed to show that this relation is not as definite as at first assumed. In the tables given by them guinea pigs receiving 0.01 c. c. reacted severely after 14, 17, and 155 days; others, receiving 1 c. c., after 14, 17, and 155 days; and, again, another series sensitized with 8 c. c. reacted severely after similar intervals. All of these series reacted but mildly after 245 days, showing apparently that the anaphylaxis, contrary to general belief, does not last so much longer after the larger than after the smaller sensitizing doses.

⁵⁷ Otto. *Loc. cit.* See also in "Kolle u. Wassermann Handbuch," *Ergänzungsband II*, p. 241.

⁵⁸ Rosenau and Anderson. *U. S. Pub. Health and M. H. S. Hyg. Lab. Bull.* 45, 1908.

These experiments, however, as well as similar ones by other workers, have shown that, once sensitized, animals may remain so for very extensive periods. In the work of Rosenau and Anderson⁵⁹ the limit of horse serum sensitization was 245 days. A few guinea pigs, sensitized with toxin-antitoxin mixtures, gave positive reactions after 732 days; more recently they have obtained a reaction after 1,096 days.⁶⁰

In properly sensitized animals the result of a sufficient dose of antigen, given at the proper time, is very often death. When the time and quantity are so chosen that instead of death there is merely a more or less severe anaphylactic shock, the animals are immediately thereafter in a refractory condition. That is, they are no longer sensitive to further injections of the antigen. This observation was made by Otto and by Rosenau and Anderson in their pioneer investigations, was confirmed by Gay and Southard, and was subsequently very thoroughly studied by Besredka and Steinhardt.⁶¹ The last-named workers named this refractory or immune condition "anti-anaphylaxis." There is obviously a great deal of both practical and theoretical significance in this fact, and methods were sought by which such an anti-anaphylactic state might be induced in sensitized animals without subjecting them to the dangers of actual shock. It was found that this could be accomplished in a number of ways. According to Besredka and Steinhardt the injection of moderate quantities of the antigen at a time just preceding the development of hypersusceptibility, in the preanaphylactic period, will render them refractory to later injections. This preventive administration, however, must be given during the later days of the anaphylactic incubation time. If given too soon after the first injection it does not prevent eventual sensitization, though it may occasionally delay its development, acting then simply as though a larger dose had been given in the first place. Thus if antigen is given by a method of introduction and in a quantity which would justify us in expecting hypersusceptibility to be developed at the end of 12 days, we can render the animal "antianaphylactic" by a second administration given, say, on the 8th, 9th, or 10th day. If we give it on the 2d, 3d, or 4th day after the first injection, it is very likely that sensitization will proceed nevertheless. Rosenau and Anderson have also investigated the repeated injection of antigen during the incubation time, and their results would also seem to emphasize the necessity of making the preventive injection close to the time at which hypersusceptibility may be expected. If quantities of 2 c. c. were injected 10 times

⁵⁹ Rosenau and Anderson. *U. S. Pub. Health and M. H. S. Hyg. Lab. Bull.* 50, 1909.

⁶⁰ They express the belief from this that a guinea pig may remain sensitive throughout life.

⁶¹ Besredka and Steinhardt. *Loc. cit.*

in the course of 17 days, and 15 to 17 days thereafter 6 c. c. of horse serum were given, the animals showed symptoms proving that anti-anaphylaxis was but partial. If amounts of 0.001 c. c. were given 5 times in a period of 8 days, and the animals were tested 23 days later, death often ensued. It is also possible, as a number of investigators have shown, to produce the antianaphylactic state by the injection of sublethal doses, even after the time has set in at which the animals are hypersusceptible. This method can be carried out successfully according to Besredka by injecting very small amounts into the brain ($1/50$ to $1/400$ of a cubic centimeter). Within a few hours after such an injection the animals may withstand an otherwise fatal dose with slight or no symptoms. Although it is generally stated that intraperitoneal injections, carried out after hypersusceptibility has set in, must be of considerable quantity (large enough to cause symptoms) in order to induce antianaphylaxis, Besredka⁶² states, in a recent résumé, that $1/50$ to $1/100$ cubic centimeter injected intraperitoneally and giving "practically no symptoms" in a sensitized guinea pig, after the anaphylactic state has set in, may render the animal entirely refractory after 5 hours.

On the other hand, Rosenau and Anderson,⁶³ working with *subcutaneous* injection, obtained results which differ considerably from those of Besredka. They sensitized a series of guinea pigs with mixtures of toxin and antitoxin, and 48 days later, at a time when the animals were hypersusceptible, gave 20 subcutaneous injections of 0.001 c. c. daily. Two days after the last injection, 0.2 c. c. of horse serum was given intracerebrally, and all of the animals showed symptoms, and many of them died. They conclude, therefore, that the repeated injection of small amounts of antigen into sensitized animals has no appreciable effect. Besredka, also, has shown by experiment that the introduction of large amounts of antigen into the previously cleansed rectum of sensitive animals is entirely without danger and will produce an antianaphylaxis, which becomes evident after 12 hours. This is probably dependent upon the very slow penetration of small amounts of antigen into the circulation from the gut, and has, therefore, an effect similar to the repeated injection of small amounts directly, or the very slow and gradual method of intravenous injection advocated by Friedberger for the prevention of serum sickness in man. This phase of the subject is considered in greater detail in a subsequent discussion of serum sickness.

Antianaphylaxis produced in this way is specific,⁶⁴ although, as

⁶² Besredka in "Kraus u. Levaditi Handbuch," *Ergänzungsband I*.

⁶³ Rosenau and Anderson. *Loc. cit.*, *U. S. Pub. Health and M. H. S. Hyg. Lab. Bull.* 45, 1908.

⁶⁴ Pfeiffer has recorded an exception to this in that he claims to have rendered a horse-serum-sensitive animal refractory by an injection of swine serum.

we shall see, there are other methods by which it is claimed that a nonspecific antianaphylaxis can be produced. One of these consists in the injection of anaphylactic animals with peptone. The problem of peptone poisoning and its relation to anaphylaxis will receive separate consideration.

Banzhaf and Steinhardt⁶⁵ have reported that 0.5 gram of lecithin given to sensitized guinea pigs protects them against second injection. Rosenau and Anderson⁶⁶ have failed to confirm this.

The above methods of rendering animals antianaphylactic, apart from the bearing they may have on purely therapeutic possibilities, serve to throw much light upon the mechanism of the reaction within the animal body. It is of great interest for the understanding of the physiological conditions underlying anaphylaxis also to consider briefly the influence upon anaphylactic shock which may be exerted by certain drugs. The preventive influence of atropin we have already mentioned in connection with the work of Auer and Lewis. Besredka, who, as we shall see, attributes the major part of anaphylactic manifestations to reactions proceeding from the central nervous symptom, claims to have succeeded in injecting ordinarily fatal doses of antigen without harm into guinea pigs previously anesthetized with ether. Banzhaf and Famulener⁶⁷ have similarly prevented shock by large doses of chloral hydrate. Rosenau and Anderson⁶⁸ could not prevent death with ether, and in similar investigations with urethane, paraldehyd, chloral hydrate, and magnesium sulphate, concluded that none of these drugs has any noticeable effect upon anaphylactic shock in guinea pigs.

Up to the present time we have confined ourselves to the description of the basic anaphylactic experiment, which is spoken of as "active sensitization" in analogy to the expression "active immunization," since, like the latter, it conveys the conception that the state of hypersusceptibility (like the immunity in active immunization) is here acquired by reason of physiological changes directly induced in the treated animal in reaction to the first injection of the foreign antigen. There is another method of inducing hypersusceptibility which, in continuance of the analogy to immunization, is spoken of as "passive anaphylaxis," since it consists in transferring the hypersusceptible condition to a perfectly normal animal by injecting into it serum from an actively sensitized one. The normal animal is thus merely the passive recipient of the reaction bodies produced in the sensitive animal by preliminary treatment.

⁶⁵ Banzhaf and Steinhardt. *Proc. Soc. Exp. Biol. and Med.*, Vol. 7, 1910.

⁶⁶ Rosenau and Anderson. *Hyg. Lab. Bull.* 64, 1910.

⁶⁷ Banzhaf and Famulener. *Studies N. Y. Dep. Health Res. Lab.*, 1908, p. 107.

⁶⁸ Rosenau and Anderson. *Jour. Med. Res.*, Vol. 21, N. S., 16, 1909.

That such a passive transference of anaphylaxis is possible was shown by a number of investigators almost simultaneously and M. Nicolle,⁶⁹ in February, 1907, published a study on the phenomenon of Arthus in which he showed that, if the serum of a hypersusceptible rabbit (sensitized with horse serum) was injected into a normal rabbit, the recipient was rendered sensitive, so that the subcutaneous injection of horse serum, made 24 hours later, produced typical infiltrations. Richet⁷⁰ soon after this succeeded in transferring hypersusceptibility toward mytilocongestin (a mussel poison) from a sensitized to a normal dog by injecting considerable amounts of the blood from the former into the latter. In this case, too, the hypersusceptibility of the second dog did not appear until one or two days after injection of the blood. At almost the same time Otto⁷¹ and Friedemann⁷² independently succeeded in transferring serum anaphylaxis from hypersusceptible to normal guinea pigs in a similar way. Experiments of Gay and Southard,⁷³ published during the same year, may possibly be also interpreted as instances of passive anaphylaxis, although their experimental procedure renders this doubtful, even in their own opinions. They injected 0.1 c. c. of serum from both sensitive and refractory guinea pigs into normal animals and followed this, after 10 days, with injections of antigen. The fact that such animals reacted may be interpreted in a number of ways. They themselves regarded the hypersusceptibility which the injected animals developed as a "purely active one," and it is more than likely that this was the case, the recipient animals being actively sensitized by traces of antigen remaining unassimilated in the blood of the actively sensitized donors. In the following year (1908) the facts of passive sensitization were rapidly confirmed and extended by Besredka,⁷⁴ Lewis,⁷⁵ and others,⁷⁶ and information of the greatest value for the comprehension of the anaphylactic reaction was obtained.

Otto showed that passive sensitization could be carried out with the serum of an actively sensitized animal 8 days after the antigen injection, at a period when this animal itself had not yet become hypersusceptible. He also showed that the passive transfer of anaphylaxis need not be confined to animals of the same species, but that guinea pigs could be rendered passively anaphylactic with the blood serum of sensitized rabbits. From the work of Gay and Southard,⁷⁷

⁶⁹ M. Nicolle. *Ann. de l'Inst. Past.*, Vol. 21, 1907.

⁷⁰ Richet. *Ann. de l'Inst. Past.*, Vol. 21, 1907.

⁷¹ Otto. *Münch. med. Woch.*, No. 34, 1907.

⁷² Friedemann. *Münch. med. Woch.*, No. 49, 1907.

⁷³ Gay and Southard. *Jour. Med. Res.*, Vol. 16, 1907.

⁷⁴ Besredka. *Ann. de l'Inst. Past.*, Vol. 22, 1908.

⁷⁵ Lewis. *Jour. Exp. Med.*, Vol. 10, 1908.

⁷⁶ Kraus and Doerr. *Wien. klin. Woch.*, No. 28, 1908.

⁷⁷ Gay and Southard. *Jour. Med. Res.*, Vol. 18, 1908.

moreover, it appeared that not only by the blood of sensitive animals can anaphylaxis be transferred, but that this can also be done by injecting the blood of animals that have once been sensitive but have subsequently been rendered antianaphylactic or refractory. Analogous to this observation is the fact, observed by these authors as well as by Friedemann, that the young of antianaphylactic mothers are not refractory but hypersusceptible. This observation, unquestionably correct, since it has been confirmed by several other workers, is astonishing and contrary to expectation. It has had no inconsiderable bearing upon our theoretical understanding of anaphylaxis.

It was soon found out, too, that hypersusceptibility was conveyed not only by the sera of sensitive and of refractory animals, but that it could likewise be transferred by the precipitating sera of animals systematically immunized with a foreign protein.

This method was later employed by Doerr and Russ⁷⁸ in their quantitative studies on the relations between anaphylactic antigen and antibody. We are confronted, then, with the curious facts that animals may be passively sensitized:

(a) by the serum of a sensitized animal.

(b) by the serum of an animal not yet sensitive—in the pre-anaphylactic period (8th day, Otto).

(c) by the serum of an antianaphylactic animal.

(d) by the precipitating serum of an "immunized"⁷⁹ animal.

Lewis further showed that normal guinea pigs could be rendered hypersusceptible with the blood of congenitally sensitive animals.

Passive sensitization is carried by the blood serum purely, since, in ordinary cases, as Rosenau and Anderson have shown, the blood corpuscles and tissues of a sensitive animal do not convey the hypersusceptibility. An exception to this will be noted later when we come to discuss Bail's experiments on the passive transfer of tuberculin sensitiveness.

Passive sensitization, once established, may persist for as long as 3 or 4 weeks, though Rosenau and Anderson found that animals tested 26 days after treatment reacted but weakly. In the young of anaphylactic mothers Otto has observed positive reactions as long as 44 days after birth, though fatal results were obtained in pigs only a few days old.

Throughout the earlier investigations upon passive sensitization the curious fact recurs in the experiments of successive workers that

⁷⁸ Doerr and Russ. *Zeitschr. f. Immunitätsforschung*, Vol. 3, 1909.

⁷⁹ We must never forget that the term "immunized" as applied to animals treated with harmless protein is an analogy and not absolutely correct. Such animals, though probably capable of assimilating larger quantities of foreign injected protein than normal ones, and this more rapidly, may nevertheless be not a whit more tolerant of the antigen—sometimes even extremely sensitive and vulnerable.

a definite period must elapse between the injection of the sensitive blood and that of the antigen.

Both Friedemann and Otto found that when the sensitive serum was injected subcutaneously the best results were obtained by administration of the antigen 24 to 48 hours after this. On intraperitoneal injection of the sensitizing serum Doerr and Russ⁸⁰ obtained the best results by permitting an interval of 24 hours to elapse, and the same investigators still further shortened this period to 4 hours by injecting the sensitive serum intravenously. Beyond this, the interval could not be shortened with success. Indeed, some writers, notably Gay and Southard, have claimed that the maximum hypersusceptibility in guinea pigs treated with sensitive serum is reached only after 10 or more days, and Rosenau and Anderson, Lewis, and others have obtained results which seem to point in the same direction. However, as we have already indicated, the testing of animals so long after the injection of sensitive serum leaves us in doubt whether we are dealing with true "passive" transference of anaphylaxis or with active sensitization due to traces of antigen carried over with the serum of the sensitive animal. For the purposes of theoretical deduction, therefore, it is better to ignore these cases and consider chiefly passive transference in which reactions are obtained within 24 hours or less after the injection of the anaphylactic serum—an interval so short that active sensitization can hardly be considered as a reasonable possibility.

The important point, in this connection, is the fact that in most of the earlier investigations it was found that between the administration of sensitive serum and of antigen a definite interval, however short, was invariably necessary.⁸¹

From these observations the natural deduction was made that the anaphylactic symptoms were the result of cellular occurrences, and that the antigen could act only after the sensitizing substance (however conceived) had become attached to certain cells, probably to those of the central nervous system. It was thought that a meeting of antigen and the sensitizing substance in the circulation would result in no reaction; that, in other words, the effective reaction

⁸⁰ Doerr and Russ. *Zeitschr. f. Immunitätsforschung*, Vol. 3, p. 181, 1909.

⁸¹ An exception to this, contradicting the then prevailing opinion, were the researches of Weill-Hallé and Lemaire (*C. R. de la Soc. de Biol.*, Vol. 65, July, 1908, p. 141), who showed that, under certain conditions, guinea pigs would react with typical, often fatal, anaphylaxis if injected simultaneously with the serum of sensitized rabbits and the antigen horse serum. According to them, the success of such experiments depended entirely upon the condition of the sensitive serum—that is, the time at which the rabbits treated with horse serum were bled. These experiments, we shall see, were later confirmed. We record them, though important, in a footnote, since we wish at present to emphasize the reasoning which led to the assumption of a cellular participation in the reaction.

was not a direct, but an indirect, one after the anaphylactic "antibody" of the sensitive serum had become bound to the cells. It will be necessary to recur to this problem when we discuss the various theories of anaphylaxis. For the present it will suffice to state that the problem has been greatly complicated because of subsequent work which, in agreement with Weill-Hallé and Lemaire, has shown that an interval is not always necessary. Richet⁸² brought evidence of this in experiments with crepitin in 1909. It will be interesting to note that he spoke of his experiments as "réaction anaphylactique *in vitro*." He sensitized a dog to crepitin, then bled him during the hypersusceptible period, mixed the serum with a harmless dose of crepitin, and injected the mixture into a normal dog. Violent anaphylaxis resulted almost immediately.

At about the same time Friedemann⁸³ published his very important studies on the mechanism of anaphylaxis in rabbits. He found that passive sensitization in these animals, in contrast to the work of others upon guinea pigs, was best obtained by the simultaneous intravenous injection of antigen and anaphylactic serum. If the injection of the sensitive serum preceded that of the antigen by as much as 24 hours, the reaction became indistinct (undeutlich), and Friedemann concluded that here, at least, there could not be assumed the necessity of preliminary sensitization of the body cells by the anaphylactic serum, as is the case in guinea pig anaphylaxis. The anaphylactic poison, whatever it may be, Friedemann concludes, is, in rabbits at least, formed in the circulating blood. In the same communication Friedemann showed that a poisonous substance, which would give rise to the symptoms of anaphylaxis, could be produced by allowing fresh alexin or complement to act upon sensitized red blood cells. These results, of the utmost importance to our knowledge of anaphylaxis, will be considered in greater detail in a succeeding section.

His observations upon rabbits, together with Weill-Hallé and Lemaire's work on guinea pigs, largely contradicting the views concerning the necessity of an interval between the two injections in passive anaphylaxis, left the problem in considerable confusion, and work especially aimed at this point was undertaken by Biedl and Kraus⁸⁴ and others. The outcome of this was to show that apparently even in guinea pigs it was possible to produce anaphylaxis in passively sensitized animals without allowing an interval between injections of the two necessary factors. Biedl and Kraus found that shock could ensue in guinea pigs even when the two substances, sensitive serum and antigen, were mixed *in vitro*, and that animals so treated were subsequently anti-anaphylactic. They, too, record

⁸² Richet. *C. R. de la Soc. Biol.*, Vol. 66, June, 1909.

⁸³ Friedemann. *Zeitschr. f. Immunitätsforschung*, Vol. 2, June, 1909.

⁸⁴ Biedl and Kraus. *Zeitschr. f. Immunitätsforschung*, Vol. 4, 1910.

that successful accomplishment of such experiments depends largely upon the properties of the sensitive sera, for they say:

"We, too, often had sera which would sensitize guinea pigs only after an interval of 24 hours and produced no effects when injected simultaneously with the antigen." It is in this difficulty probably that we must seek the discrepancy between these later results and those obtained in the earlier investigations, and it is upon this point that many of the later controversies concerning the intravascular or cellular localization of the anaphylactic reaction have turned. It will find further consideration in a later paragraph.

CHAPTER XVI

ANAPHYLAXIS (*Cont.*)

FURTHER DEVELOPMENT AND THEORETICAL CONSIDERATIONS

WE have now briefly considered some of the fundamental facts which the earlier investigations upon anaphylaxis have revealed and, although there are still many important observations to record, the material so far outlined will serve as a basis for a brief consideration of the views that have been formulated concerning the mechanism of anaphylactic phenomena.

It is clear that the chapter of anaphylaxis is hardly more than well begun. In the earlier stages of the investigations into this problem many opinions were advanced which served the valuable functions of working hypotheses, but were quickly altered, trimmed, or expanded as new and incompatible facts were revealed in astonishingly rapid succession. The final solution is probably still far beyond our present horizon, but the recent knowledge of the toxic derivatives of proteins, "the anaphylatoxins," foreshadowed by the work of Vaughan and his associates, more definitely determined by Friedemann and especially by Friedberger, has furnished hope that we are not only on the right path toward understanding anaphylaxis, but has given us a new clue to the correlation of this condition with immunity.

It will greatly facilitate exposition of the various theories which have been advanced if we bear in mind that, although there have been many discrepancies on minor phases, the differences of opinion have centered upon the cardinal points.

These are: 1. Is the anaphylactic phenomenon a true antigen-antibody reaction in which the sensitizing injection gives rise to the formation of a specific antibody with which it reacts on second injection? 2. Is sensitization the result of effects exerted upon the tissue cells, which participate directly in the reaction, or may the reaction take place entirely in the circulation, the tissue cells being affected secondarily only?

Upon these two questions we can logically classify theories of anaphylaxis.

Among the earliest definitely stated theories is that of Gay and

Southard.¹ These workers are emphatic in denying that anaphylaxis has the nature of an antigen-antibody reaction. Their views are summarized in the following as nearly as is feasible in their own words:

Increased susceptibility in the sensitized animal is due to the continued presence in the circulation of an unneutralized element of the antigen (in their case horse serum), which they call "anaphylactin," which acts as an irritant or stimulant to the body cells, and, in some way, causes them to assimilate over rapidly certain other elements of horse serum. These assimilated or toxic elements are the same as those eliminated without producing intoxication during the incubation period following the first dose. This overassimilation after anaphylaxis is the cause of the intoxication.

Gay and Southard find much support for their contentions in the results of experiments done with the so-called "passive" transfer of hypersusceptibility. As mentioned above, hypersusceptibility may be transferred to a normal animal with the blood serum not only of a sensitive animal, but even more surely and effectually with that of a refractory, or "antianaphylactic," animal. They believe that such transfer is not "passive" but "active" sensitization, being accomplished by the transfer of "anaphylactin" to the normal animal. The refractory animal has received more horse serum than the merely sensitive one, since antianaphylaxis is produced by massive injections. Therefore its blood contains more anaphylactin and is consequently more active in transferring sensitiveness. The fact that a considerable incubation time is necessary in active sensitization they attribute to the gradual action of the anaphylactin.

In passive sensitization, therefore, they assumed a similar gradual irritation of the vulnerable cells by the anaphylactin and, as we have seen, obtained their reactions in animals so treated, usually 10 to 14 days after the sensitive serum had been given. This conception of the mechanism of passive anaphylaxis was, of course, rendered unlikely by the demonstrations by Friedemann, Otto, and others that shock could be elicited in passively sensitized animals within 24 hours or less after transfer of the anaphylactic serum.

To this, however, Gay and Southard² answer by implying that this greater speed of development of sensitiveness in the experiments of Otto is due to the larger doses used by him. They say "if the doses are sufficient it (transmitted sensitiveness) may be shown in a single day (Otto)." However, it is very likely that the sensitiveness noted by them in animals two weeks after the transference of anaphylactic serum was actually positive sensitization with antigen rests, entirely comparable to the usual "Theobald Smith" phenomenon.

¹ Gay and Southard. *Jour. Med. Res.*, Vol. 16, 1907; Vol. 18, 1908; Vol. 19, 1908.

² Gay and Southard. *Jour. Med. Res.*, p. 427, 1908.

Gay and Southard's definite objections to the possibility of an antigen-antibody reaction are found in the following arguments based on experimental observations:

1. Sensibility persists for a long time, antibodies disappear rapidly.

2. In the serum of animals sensitive to horse serum antibodies to this serum are not demonstrable by complement fixation.

3. Although sensitiveness can be transferred to a normal animal, nevertheless a definite period of incubation must elapse before the recipient becomes sensitive.

To the first of these arguments Besredka³ objects by saying that, while it is true that sensitiveness persists for a long time, the power to transmit anaphylaxis passively disappears rapidly as Otto, Richet, and others have shown.

The second contention is contradicted by the work of Nicolle and Abt.⁴ But since these workers made their observations upon rabbits their experiments do not necessarily contradict those of Gay and Southard. This point at best is a difficult one to determine, especially as recent investigations have shown us that under certain circumstances antigen and antibody may be found side by side in the same serum without uniting and without therefore fixing alexin or complement.

The point of their third argument has been discussed above.

It is clear that Gay and Southard separate distinctly the substance in the antigen which sensitizes from that which exerts the toxic action on second injection.

Another theory which is based on such a separation of a sensitizing and shock-producing element in the original antigen is that of Besredka.

Besredka⁵ assumes that in the injected antigen (serum) there are two separate substances. One of these, the *sensibilisinogen*, induces, during the time of incubation, a specific antibody (*sensibilisin*). This antibody remains in part attached to tissue cells and in part circulates freely in the blood. The other substance in the antigen he calls "*antisensibilisin*." This, at the second injection, reacts with the sensibilisin and anaphylactic shock results. The nature of the symptoms is explained by the fact that the antibody or sensibilisin is attached to cells of the central nervous system, and shock can result only when such attachment is present. Thus, in passive transference of sensitization, the property of hypersusceptibility is bestowed upon the normal animal by the sensibilisin or antibody present in the circulating blood, but the significance of this

³ Besredka. *Bull. de l'Inst. Past.*, 6, 1908, p. 826.

⁴ Nicolle and Abt. *Ann. de l'Inst. Past.*, Vol. 22, p. 132, 1908.

⁵ Besredka. *Loc. cit.*

body for anaphylaxis is not in evidence until a connection with the central nervous system has been established.

There is much in Besredka's theory which is at variance with prevailing conceptions of biological phenomena of this category. The fact that an antigen should give rise to an antibody which reacts not with the substance that induced it, but with a third body, is quite out of keeping with experience.

However, it is clear that in both theories, that of Gay and Southard, as well as that of Besredka, the cardinal point is this separation in the antigen of two substances, a sensitizing and a toxic or shock-producing, and, since this forms the chief argument against an antigen-antibody conception of anaphylaxis, it will be necessary to examine the experimental evidence on which it is based.

Gay and Adler⁶ attempted to show such a dual function of the original antigen by chemical methods. They report that, by fractional precipitation of horse serum with ammonium sulphate, the successive protein fractions obtained, as saturation is increased, are found to be less sensitizing and more toxic as more and more ammonium sulphate is added. The first fraction (euglobulins) obtained by $\frac{1}{3}$ saturation is as sensitizing as whole serum and corresponds to anaphylactin, but is nontoxic when injected into sensitive animals. The last fraction, while distinctly less sensitizing than either the whole serum or the first fraction, is at least as toxic as the whole serum.

In these experiments, therefore, we have a strong argument in favor of the separate presence in an anaphylactic antigen of two bodies, the one sensitizing and the other toxogenic. However, this assertion has not remained unchallenged.

Pick and Yamanouchi,⁷ whose extensive investigation cannot be fully reviewed here, were unable to obtain such a separation; in fact, they conclude that the same substances which sensitize are also toxic, and, working with a large variety of methods, find that both the sensitizing and toxogenic properties of proteins show no differences either in chemical condition or in resistance to chemical agents or heat.

The work of Pick and Yamanouchi, however, was done with rabbits and, therefore, as bearing on the theory of Gay and Southard, the work of Doerr and Russ⁸ is more directly to the point. These workers using guinea pigs, and both horse and beef sera, obtained results which are practically diametrically opposed to those of Gay and Adler. They found that the euglobulins, obtained by $\frac{1}{3}$ saturation with ammonium sulphate, are the most strongly sensitizing and, at the same time, the most toxic of the fractions of the sera. As

⁶ Gay and Adler. *Jour. Med. Res.*, Vol. 13, 1908.

⁷ Pick and Yamanouchi. *Zeitschr. f. Immunitätsforschung*, 1, 1909.

⁸ Doerr and Russ. *Zeitschr. f. Immunitätsforschung*, Vol. 2, 1909.

saturation with the salt is increased, the proteins which come down decrease progressively and in parallelism, both as regards the power to sensitize and the faculty of exerting toxic action on second injection. The albumin, which finally comes out on total saturation, is devoid both of sensitizing and of toxic properties. Similar results were obtained by Doerr and Russ with the precipitation of serum proteins with CO_2 .

The weight of evidence, therefore, seems to point against a chemical separation of the two functions in the antigen.

Besredka's contentions in favor of such a separation were based chiefly upon a difference in resistance to heat.

His experiments showed that the sensitizing properties of serum are not lost even if it is heated to 120°C ., while the toxogenic powers are destroyed by much lower temperatures. The results of Besredka as to the differences in thermostability between the two properties have found confirmation by Kraus and Volk⁹ and others, and there can be little doubt that the sensitizing function is extremely heat-resistant, since this has also been shown by Wells,¹⁰ Rosenau and Anderson, and many others. However, researches by Doerr and Russ,¹¹ and notably by Wells, have shown that, though not *destroyed* by high temperatures, even moderate heating markedly diminishes the sensitizing function, and that larger doses have to be given as the temperature is increased; and since the smallest quantities of antigen necessary for inducing shock at the second injection must be anywhere from 100 to 1,000 times as large as the smallest sensitizing doses, it is quite likely that a combination of such conditions might simulate an actual difference in heat resistance. In fact, this is the view expressed by Wells¹² and borne out by experiments carried out by Doerr and Russ.

Wells, too, confirms the identity of sensitizing and toxic substance by his experiments on the influence of tryptic digestion upon these properties of the antigen. He concludes that both sensitizing and intoxicating properties are attacked and slowly decrease as the coagulable protein disappears.

As to that aspect of Besredka's theory which deals with the indirect participation of the central nervous system, his arguments are based mainly on the fact that ether narcosis seemed, in his experiments, to prevent anaphylactic shock when animals were deeply anesthetized during the second injection, and also upon the regularity, severity, and speed with which anaphylactic symptoms follow injections directly into the brain. The former contention regarding narcotics cannot, by any means, be accepted as yet,

⁹ Kraus and Volk. *Zeitschr. f. Immunitätsforschung*, Vol. 3, 1909.

¹⁰ Wells. *Jour. Inf. Dis.*, Vol. 5, 1908.

¹¹ Doerr and Russ. *Loc. cit.*

¹² Wells. *Jour. Inf. Dis.*, Vol. 6, p. 521, 1909.

since Rosenau and Anderson¹³ failed to confirm it and claim that ether narcosis merely masks the symptoms but does not prevent death. If we admit the beneficial effects of ether, moreover, it may well be that this is accomplished by relaxation of the bronchial spasms, known, since Auer and Lewis, to be the cause of death in guinea pigs, and the action of ether could hardly be utilized, therefore, to argue in favor of a central localization of the anaphylactic process.

That phase of the two theories so far mentioned, therefore, which depends upon the assumption of two separate substances in the original antigen does not seem established nor even sufficiently likely to warrant the formulation of a theory upon it.

The second premise is the necessary participation of the body cell, in that the reaction cannot take place unless the cells are rendered vulnerable by preliminary alteration. In Gay and Southard's theory this is accomplished by irritation exerted by the "anaphylactin;" in Besredka's scheme this is due to the antisensibilisin which is attached to the nerve cells. In both cases a gradual preliminary preparation of the cells is necessary, a view which is still held by many observers on strong evidence, although we know from the cited experiments of Friedemann, Biedl and Kraus, and others, that anaphylaxis can be produced in a normal animal by the injection of previously mixed antigen and sensitive serum, if the experimental conditions are properly understood and observed.

All other views of the mechanism of anaphylaxis have held that, in substance, this reaction is a true antigen-antibody reaction. The injected antigen gives rise to a specific antibody. This, on second injection, unites with the first antigen and the result is anaphylactic shock. Such a point of view was held from the beginning by v. Pirquet, Rosenau and Anderson, and others, who reached this conclusion from the nature of the anaphylactic antigens, the specificity of the reaction, the incubation time, and the phenomena of passive sensitization.

The conception of cell participation, however, has also been a feature of a number of theories which have interpreted anaphylaxis from the beginning as a true antigen-antibody reaction. When we come to consider anaphylaxis-like phenomena we will have a few words to say regarding the hypersusceptibility against bacterial toxins which was noticed long before the days of anaphylaxis investigations by von Behring and his pupils. To explain this occurrence Wassermann, Kretz, and others advanced the theory of "sessile receptors."

In order to make the meaning of this term clear let us briefly review Ehrlich's opinion regarding the formation of antibodies.

¹³ Rosenau and Anderson. *Loc. cit.* and *U. S. Pub. Health and M. H. S. Hyg. Lab. Bull.* 45, p. 22, 1908.

When a foreign antigen is injected into an animal the assimilation takes place by means of its entering into relation with the body cells by becoming attached to an atom-complex or cell receptor for which the particular antigen has affinity. In consequence this atom-complex, side chain, or receptor is eliminated from usefulness and the cell is forced to produce another or others like it. In ordinary immunization overproduction results, the receptors are cast off, and, in the circulation, represent the antibodies which we have studied. Now it is conceivable that slight stimulation of the cells, insufficient to induce a very extensive receptor formation, might lead to the increase of receptors without leading to their extrusion from the cell into the circulation. The condition of the cell in consequence is one of increased receptor apparatus or affinity for the particular antigen, and consequently greater vulnerability, if the antigen, as in the case of the toxins, is a harmful substance. Adapting this ingenious idea to the explanation of serum anaphylaxis, Friedberger,¹⁴ in his first theory, combined the conceptions of antibody formation and cellular localization. He identified the anaphylactic reaction with the precipitins and advanced the opinion that the anaphylactic reaction was a sort of intracellular precipitin reaction.

In the light of the evidence against the histogenic conceptions of anaphylaxis which we have mentioned above, and especially because of his own discoveries upon the anaphylactic poisons, Friedberger has abandoned this view, and no more need be said about it at present. However, his identification of the precipitins with the anaphylactic antibody is of great interest in that it stimulated much careful analytical work on the antibodies present in anaphylactic sera.¹⁵

Thus, the assumption that anaphylaxis was in truth the result of the union of an antigen with its specific antibody gained much support when Doerr and Russ¹⁸ succeeded in applying quantitative methods to the study of the anaphylactic antibody. Their methods consisted in producing precipitating sera in rabbits. With these they then passively sensitized guinea pigs, subsequently testing them with the antigen 24 hours later. To arrive at quantitative results they developed two reliable methods. These consisted in: 1. Intraperi-

¹⁴ Friedberger. *Zeitschr. f. Immunitätsforschung*, Vol. 2, 1909, p. 208.

¹⁵ The idea of identifying the anaphylactic antibody with the precipitins, indeed, had been advanced before this by Hamburger and Moro,¹⁶ who believed that the first injection gave rise to precipitins—these with the antigen formed precipitates which then caused embolic obstructions. Such a purely mechanical theory soon had to be abandoned, however, because the injection of massive reëmulsi-fied precipitates did not seem to cause illness in animals and precipitins could not be demonstrated in the sera of sensitive animals.¹⁷

¹⁶ Hamburger and Moro. *Wien. klin. Woch.*, Vol. 16, No. 15, 1903.

¹⁷ Marfan and LePlay. Cited from Levaditi.

¹⁸ Doerr and Russ. *Zeitschr. f. Immunitätsforschung*, Vol. 3, pp. 181 and 706, 1909.

toneal sensitization of guinea pigs with constant quantities of titrated precipitating serum. Twenty-four hours later intravenous test with diminishing amounts of specific antigen. 2. Intraperitoneal sensitization with diminishing quantities of the titrated precipitating serum, and 24 hours later intravenous tests with constant amounts of antigen.

In this way they showed that there was a direct relationship between the power of a serum to convey anaphylaxis passively and its contents of precipitins. We may elucidate this by an example from their work. They possessed a rabbit serum which gave precipitation with sheep, goat, beef, pig, human, and horse sera, but not with chicken serum. The precipitation titre of this serum for the sera mentioned varied from 1 in 20,000 in the case of sheep and goat sera, to 1 in 100 in the cases of the human and horse sera. When guinea pigs were injected intraperitoneally with 1 c. c. of this serum, and after 24 hours were intravenously injected with the various sera mentioned above, in decreasing quantities, the sera which were precipitated in the highest dilutions gave anaphylactic shock in the smallest quantities. Those sera for which no precipitin or little had been present in the antiserum gave little or no reaction by this method even where considerable quantities were used. Thus in animals prepared by 1 c. c. of this antiserum, the sheep serum (precipitated in dilutions of 1 in 20,000) caused death when injected in doses of 0.006 c. c., whereas horse serum (which was precipitated only in concentration of 1 to 100) gave slight symptoms only when 2 c. c. were employed for reinjection and chicken serum (non-precipitable by the antiserum) gave no reaction in similar doses.

In this, then, we have a definite quantitative analysis which proves that the power to sensitize passively is in direct relation to the antibodies against the protein present in the sensitizing serum. Whether or not this means the precipitins particularly we will consider in a later section.

We are now prepared to follow individually the development of those theories in which the anaphylactic mechanism was looked upon purely as the result of the union of an antigen with its antibody.

The conception which gradually grew out of the antigen-antibody mechanism of anaphylaxis was the following: When a specific antigen meets its antibody the reaction between them gives rise to a toxic product, and this causes the characteristic symptoms. A similar idea, it will be remembered, is found in the original endotoxin theory of Pfeiffer. According to this, the action of the specific lysin liberated from bacteria a preformed poison, the endotoxin. In 1902 Weichhardt,¹⁹ bearing this conception in mind, subjected syncytial protein of rabbit placenta to the action of specific antisera and obtained substances toxic for normal rabbits.

¹⁹ Weichhardt. *Deutsche med. Woch.*, 1902, p. 624.

This work was done long before the days of anaphylaxis studies, and the results were interpreted in keeping with Pfeiffer's theory. However, as Weichhardt himself now claims, it is not unlikely that he was dealing with a phenomenon analogous to the ones we are discussing. A similar opinion of the production of toxic substances by specific cytolysis was expressed by Wolff-Eisner²⁰ in 1904.

Probably the most important of the earlier investigations along these lines, at least in its direct bearing on anaphylaxis, was the work of Vaughan and Wheeler,²¹ published in 1907.

In its general significance this work ranks among the most important contributions to our understanding of hypersusceptibility.²² Their conception of anaphylaxis takes root in the earlier investigations of Vaughan²³ and his pupils upon the extraction of a poisonous group from the protein molecule.

Vaughan and Wheeler²⁴ ²⁵ believe that the sensitizing and the toxogenic properties of the anaphylactic antigens are in truth contained within the self-same proteid molecule; but can be chemically separated from each other. They have been able to split egg albumen and other proteids by treatment with absolute alcohol (containing 2 per cent. NaOH) into 2 fractions—a toxic alcohol-soluble and a non-toxic alcohol-insoluble one. The former fraction gave protein reactions, and they regard it as a true protein—while Wells,²⁶ considering the hydrolytic nature of the cleavage resorted to, considers this fraction as possibly a soluble peptone or polypeptid (the positive protein reactions being possibly due to amino acids). The non-alcohol-soluble, non-toxic fraction also gives proteid reactions. Injections into guinea pigs of the toxic fraction produce symptoms not unlike anaphylaxis—but do not sensitize against protein. The alcohol-soluble portion is non-toxic and sensitizes against protein in doses of 0.001 to 0.005 gm.

Based on these results, their views of mechanism of anaphylaxis are as follows: At the first injection a slow lysis (cleavage) of the injected protein gradually liberates a fraction, corresponding to the alcohol-insoluble substance—and this by its antigenic action gives rise to the formation, in excess, of an enzyme (lysin), which on re-injection brings about the rapid cleavage of the injected protein—

²⁰ Wolff-Eisner. *Centralbl. f. Bakt.*, Vol. 37, 1904.

²¹ Vaughan and Wheeler. *Jour. Inf. Dis.*, Vol. 4, 1907.

²² This work also contains the germ of the more recent ideas upon the nature of toxemia in infectious disease, advanced more particularly by Friedberger. This will be considered in detail in the next chapter.

²³ Vaughan. *Transact. Ass'n Am. Phys.*, Vol. 16, 1901; *Jour. A. M. A.*, Vol. 36, 1901; *Am. Med.*, 1901; *Jour. A. M. A.*, Vol. 43, 1904.

²⁴ V. C. Vaughan, Jr. *Jour. A. M. A.*, Vol. 44, 1905, p. 1340.

²⁵ V. C. Vaughan. *Boston Med. and Surg. Jour.*, Vol. 155, 1906.

²⁶ Wells. *Jour. Inf. Dis.*, Vol. 5, 1908.

with an explosive liberation of the toxic fraction and consequent symptoms.²⁷

Nicolle believes that the injection of a protein into an animal induces the production in the subject of antibodies. These are preëminently two—albuminolysins, which cause its cleavage, and albuminocoagulins or precipitins, which coagulate and prevent the action of the lysin. At the time at which an animal is hypersusceptible or anaphylactic there has been a production of albuminolysins which cause cleavage of the protein, with the rapid liberation of toxic substances; but the albuminocoagulins or precipitins have not yet adequately developed. In a refractory animal the neutralizing action of the albuminoprecipitins prevents the harm which the lytic action might otherwise accomplish. The relative amounts of these two antibodies present in the circulation of the animal at any particular time determine whether the animal is anaphylactic or refractory or immune. This theory assumes arbitrarily the protective nature of precipitation, an idea which has no foundation in experiment and, in fact, is rendered extremely unlikely by more recent developments of our knowledge of the precipitating antibodies.

Given, then, a reasonable hypothesis in which anaphylaxis is associated with the cleavage of protein by lysis, given, in other words, an antigen-antibody conception, it is but natural that experimenters should ask themselves: What is the relation of the alexin to this cleavage? For in all known lytic reactions, of course, the union of antigen and antibody leads to the absorption of alexin, by means of which, then, the lysis takes place. This problem suggested itself to a number of the earlier investigators who attempted to approach it by determining whether or not the sera of sensitive animals, added to antigen, would fix alexin. Gay and Southard, Sleeswijk,²⁹ and others obtained negative results, while Nicolle and Abt,³⁰ and Doerr and Russ³¹ obtained positive results. As far as this particular method is concerned, therefore, no conclusions can be drawn. Sleeswijk, however, has approached the question in another way and examined whether or not there is a diminution of alexin in the blood of an animal immediately after anaphylactic shock. He found that this was indeed a regular occurrence, and his results have been confirmed by Friedberger and Hartoch³² and a number of others.

It was shown by these workers that, both in active and passive anaphylaxis in rabbits and dogs, as well as in guinea pigs, there is a definite and considerable diminution of complement immediately after anaphylactic shock.

²⁷ For the sake of completeness it is well also to mention Nicolle's²⁸ theory, which, though attractive, is not borne out by recent knowledge concerning the nature of precipitins.

²⁸ Nicolle. *Ann. de l'Inst. Past.*, Vol. 22, 1908.

²⁹ Sleeswijk. *Zeitschr. f. Immunitätsforschung*, Vol. 2, 1909.

³⁰ Nicolle and Abt. *Ann. de l'Inst. Past.*, Vol. 22, 1908.

³¹ Doerr and Russ. *Zeitschr. f. Immunitätsforschung*, Vol. 3, 1909.

³² Friedberger and Hartoch. *Zeitschr. f. Immunitätsforschung*, Vol. 3, 1909.

The question now arises: What is the significance of this diminution of alexin? Do the animals die because of a sudden loss of circulating, physiologically necessary alexin, or does the alexin take an *active* part in producing the conditions which cause death?

Either of these possibilities might follow from the mere fact of alexin diminution, but the former—the possibility that complement depletion is the cause of death—was ruled out by Friedberger and Hartoch.³³ They showed that, by supplying fresh complement to sensitive animals at the time of reinjection, shock cannot be prevented. They now proceeded to demonstrate the *active* participation of complement in the production of anaphylaxis. They did this in an ingenious way which depended on utilization of the fact observed by Nolf,³⁴ Hektoen and Ruediger,³⁵ and others that hypertonic salt solution (1.5-2 per cent.) will prevent the combination of complement with its sensitized cells. By slowly injecting into sensitized guinea pigs 0.3 cubic centimeter of concentrated NaCl solution just before the injection of antigen they were able to markedly diminish anaphylactic shock—saving animals from injections which invariably killed the controls.

An extremely ingenious demonstration of the important rôle played by complement in anaphylaxis has recently been furnished by Loeffler. Loeffler,³⁶ using guinea pigs sensitized with horse serum, completely depleted their complement by injecting intraperitoneally considerable quantities of sensitized sheep corpuscles. Tested by injection of horse serum one hour later no anaphylaxis occurred, while controls regularly succumbed.³⁷

It was thus established with as much accuracy as the peculiar experimental difficulties of the problem permitted that the complement or alexin played an important active part in the production of anaphylaxis, and the next logical step was to attempt to produce the anaphylactic poison by the action of alexin upon an antigen-antibody complex *in vitro*. This was first done, with direct reference to anaphylaxis, by Ulrich Friedemann.³⁸ Friedemann chose as his antigen-antibody complex the sensitized red blood cell after he had demonstrated by preliminary experiment that the basic anaphylactic experiment could be carried out in rabbits with washed beef corpuscles. He found that if 3 c. c. of such corpuscles were injected into rabbits and the injection repeated after 3 weeks anaphylactic symptoms were regularly elicited. He then allowed alexin to act upon sensitized beef blood *in vitro*, interrupted the action by cool-

³³ Friedberger and Hartoch. *Loc. cit.*

³⁴ Nolf. *Ann. de l'Inst. Past.*, 1900.

³⁵ Hektoen and Ruediger. *Jour. Inf. Dis.*, Vol. 1, 1904.

³⁶ Loeffler. *Zeitschr. f. Immunitätsforsch.*, 8, 1910.

³⁷ For additional evidence pointing in the same direction see also Uhlenhuth and Haendel, *Zeitschr. f. Immunitätsforsch.*, Vol. 3, 1909.

³⁸ Ulrich Friedemann. *Zeitschr. f. Immunitätsforsch.*, Vol. 2, 1909.

ing at a time just preceding the occurrence of hemolysis (to exclude the supposed toxic action of hemoglobin), and injected the supernatant fluid of such mixtures into normal rabbits. The result was marked illness resembling anaphylaxis, and Friedemann thus had succeeded in producing the anaphylactic poison *in vitro* under conditions as nearly as possible similar to those occurring in the circulation of the anaphylactic rabbit. In the conclusions drawn from his experiments he expresses the opinion that the poisons were not preformed in the red blood cells, but were formed by the proteolysis exerted by "amboceptor" and complement. In this statement he sets down the basic conception of the production of anaphylactic poisons now generally held.

Friedemann, then, in attempts to apply the same methods to the study of serum anaphylaxis, attempted to produce similar poisons by the action of rabbit alexin upon the washed precipitates formed by mixtures of antigen and precipitating sera. In this he failed—probably because of his choice of rabbits as subjects for experiment. Where he had failed, however, Friedberger³⁹ succeeded by using guinea pigs. Doerr and Russ⁴⁰ had previously shown that feeble symptoms of shock could be produced by the injection of serum precipitates into normal guinea pigs. With this additional evidence in favor of his reasoning, Friedberger proceeded as follows:

One c. c. of a rabbit serum which precipitated sheep serum in a dilution of 1 to 10,000 was mixed with 30 c. c. of a 1 to 50 sheep serum dilution. This was kept one hour at 37.5° C. and over night in the ice-chest, when a heavy flocculent precipitate had formed. This precipitate was washed to remove all traces of serum, and to it were added 2 c. c. of fresh normal guinea pig serum—as complement. This was again allowed to stand for 12 hours and then the supernatant fluid was injected into a guinea pig intravenously. In most cases the pigs so treated showed marked symptoms soon after the injection and died within a few hours.

Friedberger concludes, therefore, that anaphylactic shock is a true intoxication due to a poison produced from the products of a precipitin-precipitinogen reaction by the action of a complement; he speaks of the formed poison as *anaphylatoxin*. The experiment just outlined, moreover, seems to show, contrary to Friedberger's first ideas, that the entire reaction may go on under certain circumstances in the blood stream without intervention of sessile precipitins upon the cells.

We have, thus, in the cited work of Friedberger the culmination of a long series of investigations—the end result being the conclusion

³⁹ Friedberger. *Berl. klin. Woch.*, 32 and 42, 1910; also *Zeitschr. f. Immunitätsforsch.*, Vol. 4, 1910.

⁴⁰ Doerr and Russ. *Zeitschr. f. Immunitätsforsch.*, Vol. 3, p. 181, 1909.

that in all probability—at least as far as experimental ingenuity has permitted us to penetrate into this very difficult problem up to the present time—the phenomenon of anaphylaxis must be regarded as an acute intoxication, the poison which calls it forth being the result of the union of an antigen and its antibody, the complex being subsequently subjected to proteolysis by the action of alexin or complement. The experimental extension of this conception to the phenomena of bacterial anaphylaxis has promised to exert such an important influence upon our conceptions of infectious disease that we will take up these investigations in a separate section. Although it seems proved with reasonable certainty, however, that the above mechanism accounts for anaphylactic shock as produced in the ordinary experiment, there are still a number of important questions which await further solution. Among these is primarily the question as to whether or not we are justified in excluding finally the possible primary participation of the body cell in all cases of anaphylaxis and the problem of the identification of the anaphylactic reaction body with any of the known antibodies.

Many of the arguments we have cited—notably those of Friedemann and Friedberger—seem to exclude the participation of the cells and tissues in the anaphylactic reaction as experimentally produced, or, rather, seem to show that shock can be explained without resort to cellular participation. However, there are certain experimental data which cannot be ignored which point toward the participation of cell-changes in the condition of sensitiveness; and while it is probable that anaphylactic poisons may be sufficiently formed in the circulation to cause shock in the rough procedures which must mark even our most delicate experiments, there is in addition to this much evidence of an alteration of cellular susceptibility which influences the anaphylactic phenomena. The question is not absolutely settled to-day and it is a problem in which it is useless to speculate. We must attempt to approach it, therefore, by citing the evidence which has been advanced on both sides. Some of these data we have already discussed, in connection with passive sensitization, on page 383.

The opinion of v. Pirquet was originally that there must be a change in susceptibility in the cells to account for the various skin reactions, assuming these analogous to anaphylaxis. Experiments which seem to bear this out are those of Schultz⁴¹ upon the increased sensitiveness to serum of the excised smooth muscle of anaphylactic animals. It had been shown that smooth muscle contracts when brought in contact with serum. That of sensitized guinea pigs showed a markedly greater susceptibility in this respect. Of great importance in pointing to primary cell participation also it seems to

⁴¹ Schultz. *Jour. of Pharm. and Exp. Therap.*, 1, 1910.

us is the following ingenious experiment of Pearce and Eisenbrey.⁴² We cite their own description:

"Our procedure has been to exsanguinate under ether anesthesia a small, normal dog (A), and to transfuse this animal by Crile's method with the blood of a larger sensitized dog (B) until the blood pressure reached approximately its original level. After sufficient blood had been obtained from B to raise the pressure of A the sensitized dog was then bled to exsanguination and transfused from a third normal dog (C) until its pressure reached its previous normal level. At the proper moment the normal dog containing the blood of the sensitized dog, and the latter containing the blood of the normal dog, each received intravenously the toxic dose of horse serum. In the former a fall in pressure does not occur, and in the latter it does, thus proving that the phenomenon of anaphylaxis is due to a reaction in the fixed cells, and not either primarily or secondarily in the blood."

Experiments similar in significance to those of Schultz have been carried out by Dale,⁴³ who used as his indicator of cellular susceptibility the uterine muscle of virgin guinea pigs. These experiments have been confirmed by Weil.⁴⁴ Coca,⁴⁵ too, has added to the evidence in favor of the cellular localization by experiments in which he practically repeated in guinea pigs the observations made by Pearce and Eisenbrey upon dogs, adding a number of further important points concerning the participation of the complement in anaphylaxis, which we will take up directly.

In regard to this problem of the localization of the anaphylactic mechanism, then, we are confronted with two predominant lines of evidence, each of which has seemed so well supported by experiment that it has tempted investigators into the expression of positive opinions and the formulation of theories.

On the one hand, it is unquestionable that poisons can be produced by the action of complement upon antibody-antigen complexes and that these poisons, injected into guinea pigs, produce symptoms indistinguishable from anaphylactic shock. Here there seems to be proof that anaphylaxis can be induced by agencies all of which are available in the blood stream under the conditions under which the phenomenon is observed.

On the other hand, the experiments just cited permit of no doubt

⁴² Pearce and Eisenbrey. *Transac. of Congr. of Am. Ph. and Sur.*, Vol. 8, 1910.

⁴³ Dale. *Journ. of Pharm. and Exp. Therap.*, Vol. 4, 1913.

⁴⁴ Weil. *Journ. of Med. Res.*, 27, 1913; 30, 1914.

⁴⁵ Coca. *Zeitschr. f. Imm.*, Vol. 20, 1914.

that the property of sensitiveness in anaphylactic animals may be an attribute of the cells, independent of the substances circulating in the blood. Can we reconcile these apparently opposed facts?

It would seem to us most rational to look upon the problem in the following way: The antibodies produced by the body in response to the injection of an antigen are, of course, the products of the cells, and it is likely, on the basis of experiments and data considered in another chapter, that many or all the cells of the body with which the antigen comes in contact may participate in their production. At different periods during the process of immunization the antibodies are therefore present, in varying ratio, both in the circulation and within the cells. During the earlier stages of the response to the antigen, i. e., during the period of hypersensitiveness, it is likely that the reaction changes (antibodies) are more plentiful in the cells than in the blood stream. When antigen is injected into a sensitized animal, it is likely that it goes into reaction with both the circulating and the sessile antibody. The latter is probably most important in the ordinary reaction of anaphylaxis, the former is probably of great importance in such cases as the sudden death which can be seen in highly immunized rabbits, when a fourth or fifth injection of a foreign serum is given (at a time at which the blood already strongly reacts with injected antigen). Both the cellular and the intravascular reaction probably occur in all cases, although we are inclined to believe that the cellular reaction must be taken to dominate ordinary anaphylaxis, as observed experimentally. That the intravascular reaction, however, may also have importance is testified by the experiments which we have cited (pp. 383, 384 and 398) and which cannot be ignored.

Now, granted that the reaction takes place in both the cells and the circulation, varying in relative intensity in each location according to the stage of antibody formation, and the relative concentration of the antibodies in cells and in blood stream, in how far are we justified in assuming that the complement or alexin of the circulating blood is necessary for the production of anaphylactic shock? The experiments of Friedberger and many others have shown beyond doubt that the action of complement upon antibody-antigen complexes may produce poisons, and much evidence has been gathered to show that complement is diminished in animals during shock. This constitutes reasonable ground for assuming that the complement participates at least in that phase of anaphylaxis which takes place in the blood stream. Whether or not the circulating complement acts upon those antibody-antigen complexes which are formed on the sensitive cell, is hard to decide. Experimentally it cannot be absolutely determined, since it would be quite impossible to remove all traces of complement from any cell. Moreover the complement is, after all, also a cell product, and it is more than likely that the cell disposes

over intracellular enzymes quite capable of substituting functionally for complement.

It seems necessary to add that, however one may look upon this, it does not affect the importance of the so-called "anaphylatoxins" and similar toxic protein cleavage products in the toxemia of infectious disease or in general pathology.

Now, as to the identification of the anaphylactic antibody with some one of the well-known antibodies, the assumption is that in cellular anaphylaxis (as in the corpuscle experiments of Friedemann and in the bacterial experiments of Friedberger and others) the so-called sensitizer or amboceptor is to be held responsible. This seems reasonable, and there is much evidence that seems to favor such a view.

In the case of serum anaphylaxis extensive work has been done to show a parallelism between the anaphylactic antibody and the precipitins. This we have seen principally in the experiments of Doerr and Russ, and those of Friedberger.

The problem becomes a complicated one when we attempt then to define the nature of the precipitins and their relation to the antibodies hypothetically advanced as "albuminolysins" by Gengou. Without going into this point extensively at present, it may be permitted to refer the reader to the chapters on alexin fixation and precipitins, and to reiterate the writer's⁴⁶ own opinion, which is that much reasonable evidence points to the fact that the so-called precipitins are in truth protein-sensitizers, identical in structure and function with the sensitizers or amboceptors of cytolytic processes. The fact that precipitation occurs when these antibodies are added to the homologous dissolved antigen is merely a secondary colloidal phenomenon; antigen and antibody react, forming a complex which is then amenable to the action of alexin. Being colloidal in nature, and mixed under quantitative and other conditions which favor flocculation, they precipitate. This point of view, then, identifies the so-called precipitins with the protein-sensitizers or albuminolysins first hypothetically suggested by Gengou. It leads necessarily to the conception that in cytolysis as well as proteolysis, in fact, in all reactions in which antigen is sensitized to the action of alexin, there is functionally but one variety of antibody—the sensitizer—precipitation and agglutination being incidental physical phenomena not dependent upon special antibodies as heretofore supposed.

In this sense, then, the "precipitins" or albuminolysins may be regarded as identical with the anaphylactic antibody.

That animals in whose circulation antigen and antibody are simultaneously present do not suffer from symptoms of anaphylaxis

⁴⁶ Zinsser. *Jour. Exp. Med.*, Vol. 15, 1912, and Vol. 18, 1913.

has been referred by Zinsser and Young⁴⁷ as possibly due to the action of a protective colloid which prevents the union of the two.

THE MECHANISM OF ANTI-ANAPHYLAXIS

The conditions under which animals, previously anaphylactic, may be rendered refractory or "anti-anaphylactic" have been discussed in another place. This condition is not entirely comparable with immunity since it is a purely temporary state, lasting possibly a few weeks, but after this the animals do not return to the normal condition, but gradually become again moderately hypersusceptible. (Rosenau and Anderson, Otto and others.) Thus a guinea pig which has been sensitized, then rendered anti-anaphylactic by a massive injection of antigen, may react with mild symptoms to an injection made 20 to 30 days later. Such returning sensitiveness, according to Rosenau and Anderson,⁴⁸ is usually mild, fatal reactions rarely occurring.

An entirely satisfactory theory of anti-anaphylaxis has not yet been advanced.

Besredka,⁴⁹ as we have seen, believes that the anaphylactic reaction takes place by the union of the toxic factor in the serum (anti-sensibilisin) with a specific antibody sessile upon the cells of the central nervous system. If the antigen is injected slowly or in small amount these sessile receptors are gradually united to antigen without fatal shock, and the animal is thereby rendered insensitive.

In his own words, this "desensitization" amounts to a return of the cells to their normal preanaphylactic or naturally insensitive condition. With the refutation of his theory of anaphylaxis, his theory of anti-anaphylaxis also falls to the ground, and neither of the two can be accepted as valid at present.

If we look upon anaphylaxis as a reaction taking place entirely in the circulation we may accept, with Rosenau and Anderson,⁵⁰ Friedberger, and others the explanation that anti-anaphylaxis is due to a saturation of the anaphylactic antibody with antigen. Hypersusceptibility is then subsequently reestablished because a gradual formation of circulating antibody continues, and eventually free antibody will again be present in the blood. This view is only in part satisfactory, as Friedemann⁵¹ points out. For it does not

⁴⁷ Zinsser and Young. *Jour. Exp. Med.*, Vol. 17, 1913.

⁴⁸ Rosenau and Anderson. *U. S. Pub. Health and M. H. S. Hyg. Lab. Bull.* 63, 1907.

⁴⁹ See Besredka, "Kraus u. Levaditi Handbuch, etc.," *Ergänzungsband* 1, p. 246.

⁵⁰ Rosenau and Anderson. *U. S. Pub. Health and M. H. S. Hyg. Lab. Bull.* 64, 1910.

⁵¹ Friedemann. "Frei Vereinigung f. Mikrobiol.," Berlin, 1910. Ref. *Centralbl. Bakt.* I, Vol. 47; Beiheft, p. 1, 1910.

explain the anti-anaphylaxis which Biedl and Kraus⁵² have noticed after the injection of mixtures of antigen and antibody, nor the non-specific antianaphylaxis which the same workers have observed after peptone injections. It is clear that the nature of anti-anaphylaxis remains for the present obscure, and, in view of the recent attempts to account for certain phases of infectious disease by the anaphylactic phenomena, is one of the most important problems of immunity.

Bearing upon this condition of anti-anaphylaxis is the tolerance to the anaphylactic poison which has been observed to develop in animals once or twice injected. Vaughan⁵³ has noticed this in animals injected with his toxic split products, produced by alkaline-alcohol splitting of colon bacilli. By repeated injection of the guinea pigs he showed that a tolerance was developed which protected the animal from about double the fatal dose, but the animal is not protected against larger multiples, and the condition is not an immunity in the sense in which we have used the term. Similar observations have been made by Bessau.⁵⁴ Bessau passively sensitized guinea pigs with 1 c. c. of anti-horse serum intraperitoneally, and on the following day injected them intravenously with 1 c. c. of horse serum. He gauged his dose so that the animals should suffer severe shock but survive. One or two days later he injected the amount of typhoid anaphylatoxin which was fatal for normal pigs, and found that those which had been treated as described were now able to withstand the anaphylatoxin. These experiments of Bessau would indicate that anti-anaphylaxis was to a certain extent due to tolerance of the poison, and that it was non-specific. Friedberger, together with Szymanowski, Kumagai, Odaira and Lura, later studied this problem and came to the conclusion that anti-anaphylaxis is strictly specific, depending, as Friedberger had suggested, upon the diminution of specific antibodies rather than upon tolerance to the poison. They claimed that animals that had been sensitized and then had survived the "shock" dose of homologous protein showed no tolerance for anaphylatoxin, and that animals that had been treated with the sublethal dose of anaphylatoxin were, after 24 hours, as sensitive to anaphylatoxin, however prepared, as were normal animals. Recent studies along the same lines by Zinsser and Dwyer⁵⁵ have yielded results differing from these conclusions. Working with typhoid anaphylatoxin they found that guinea pigs treated with a sublethal dose of anaphylatoxin developed a tolerance which enabled them to

⁵² Biedl and Kraus. *Zeitschr. f. Imm.*, Vol. 4, 1910.

⁵³ Vaughan. "Protein Split Products, etc.," Lea and Febiger, Philadelphia, 1913, p. 139.

⁵⁴ Bessau. *Centralbl. f. Bakt.*, Vol. 60, 1911.

⁵⁵ Zinsser and Dwyer. Reported at the meeting of Am. Ass. of Path. and Bact., Toronto, April, 1914.

resist $1\frac{1}{2}$ to 2 units of poison, the tolerance developing within three days and lasting, to a slight degree, for as long as two months. It seemed to them that animals treated with a second dose of anaphylatoxin within 24 hours after the first, if the results of this first injection have been severe, as they usually are, are still weak and generally depressed in vitality, so that a developed tolerance may be clouded by this condition. The tolerance did not seem to be strictly specific in that typhoid anaphylatoxin seemed to produce a moderate tolerance to prodigious anaphylatoxin.

It would seem, therefore, that in anti-anaphylaxis we might have two very important elements. The one, strictly specific, depends upon the depletion of antigen from the body, a true "desensitization." The other, non-specific, and probably of secondary importance since so far it has not been shown to any very powerful degree, consists of the development of tolerance by the body cells for the anaphylactic poison.

NATURE OF ANAPHYLACTIC POISON

As to the nature of the anaphylactic poison we are also to a large extent in the dark. From the experimentation upon the production of these poisons *in vitro* it appears that they are protein cleavage products. This is indirectly indicated also by metabolism experiments—such as those of Friedemann and Isaak,⁵⁶ and of Weichhardt and Schittenhelm.⁵⁷ It appeared from this work that, as measured by nitrogen output, the cleavage of foreign protein injected into specifically sensitized or immunized dogs occurred with much greater energy and speed than occurred in normal animals after first injection.

Attempts to obtain the poison by non-specific methods—that is, by purely chemical processes without the agencies of alexin and sensitizer or antibody—have been made with apparent success by Vaughan and Wheeler,⁵⁸ whose toxic, alcohol-soluble fraction (obtained by boiling egg-white in absolute alcohol containing 2 per cent. NaOH) seems to produce typical anaphylaxis in guinea pigs. This substance Vaughan and Wheeler regard as a protein, whereas Wells⁵⁹ states that it may be this, or a "soluble peptone or polypeptid, containing enough of the different aminoacids to give all the usual reactions." Weichhardt,⁶⁰ too, has obtained similar poisons by a method similar in principle to that of Vaughan and Wheeler.

⁵⁶ Friedemann and Isaak. *Zeitschr. f. exp. Path. u. Ther.*, Vol. 1, 1905.

⁵⁷ Schittenhelm u. Weichhardt. *Münch. med. Woch.*, 1910, No. 34. and 1911.

⁵⁸ Vaughan and Wheeler. *Loc. cit.*

⁵⁹ Wells. *Jour. Inf. Dis.*, Vol. 5, 1908.

⁶⁰ Weichhardt. *Centralbl. f. die ges. Phys. u. Path. des Stoffwechsels*, No. 15, 1909. Ref. "Weichhardt's Jahresbericht," 1910, p. 554.

This substance is, according to him, pharmacologically identical with his "keno toxin," or fatigue toxin, obtained in the washings from the muscles of excessively fatigued animals.

Accurate chemical definition of the anaphylactic poison has not so far been accomplished, and it is obvious that the problem is an extremely difficult one. Biedl and Kraus,^{61 62} however, have drawn a very close parallelism between anaphylactic intoxication and peptone poisoning in dogs. They have shown that peptone (0.3 gr. to the kilo.) injected into these animals gives rise to the same clinical symptoms that characterize anaphylaxis. It is accompanied also by typical fall of blood pressure, delayed coagulability of the blood, and leukopenia. Furthermore, they claim that the injection of sublethal doses of Witte peptone into serum-sensitized dogs leads to a non-specific anti-anaphylaxis. They claim a physiological identity of the Witte peptone with the anaphylactic poison.

This last observation could not be confirmed by Manwaring,⁶³ who found that dogs that had been rendered anti-anaphylactic to horse serum still reacted strongly to peptone—an observation which does not indeed weaken the contention of Biedl and Kraus as to the similarity of peptone shock to anaphylaxis, but has significance in contradicting the doubts their experiments have thrown on the specificity of anti-anaphylaxis.

Observations similar to those of Biedl and Kraus on the toxic action of peptone have been made by Arthus.⁶⁴

Biedl and Kraus have found a similar parallelism in guinea pigs in which they determined the typical bronchial spasms after peptone administration. This is in contrast to Werbitzky,⁶⁵ who found even large doses of peptone non-toxic for guinea pigs. Nevertheless, there is no question that the similarity between peptone shock and anaphylaxis is very striking and of great theoretical importance. It does not, however, bring us much nearer to a chemical understanding of the nature of the poisons since the "Witte" peptone used in these experiments is a mixture of many different substances. Brieger,⁶⁶ for instance, found toxic and non-toxic lots of Witte peptone. The toxic ones yielded on extraction a body which he calls peptotoxin. This variation in the constitution of different samples of so-called "peptone" may account for some of the conflicting results obtained in guinea pigs.⁶⁷

⁶¹ Biedl and Kraus. *Wien. klin. Woch.*, No. 11, 1909.

⁶² Also "Kraus u. Levaditi Handbuch, etc.," *Ergänzungsband* 1, p. 264.

⁶³ Manwaring. *Zeitschr. f. Immunitätsforschung*, Vol. 8, p. 589, 1911.

⁶⁴ Arthus. *C. R. de l'Acad. des Sci.*, Vol. 148.

⁶⁵ Werbitzky. *C. R. de la Soc. de Biol.*, Vol. 66, 1909.

⁶⁶ Brieger. "Die Ptomaine," 1, p. 14.

⁶⁷ For analysis of Witte peptone, see Hammarsten, "Physiological Chemistry," English translation.

PHENOMENA CLOSELY RELATED TO ANAPHYLAXIS

There are a number of well-defined phenomena of acquired hypersusceptibility or sensitiveness which, in nature, seem closely analogous to true anaphylaxis as we understand it to-day, but regarding the mechanism of which the opinions of experimenters are still to some extent at variance.

Among the most important of these is the *toxic action of normal sera* when injected into animals of another species—a phenomenon which is now generally accepted as belonging in principle to the true anaphylactic phenomena, though this opinion is of comparatively recent formulation. The subject is of sufficient theoretical and practical importance to be considered in some detail.

The older studies of phenomena belonging to this category followed closely in the footsteps of experiments on transfusion, and as early as 1666 a commission of the London Royal Philosophical Society reported deaths following transfusion, alleging intravascular coagulation as the probable cause of death.

The cause of death following injections of foreign whole blood, blood cells, and serum has, since that time, occupied the attention of many workers whose studies need not be reviewed for our present purposes. Chief among them were Magnani, Brown-Séquard, Magendie, and, more recently, Naunyn, Landois, and Ponfick.⁶⁸

The work of Landois is of special interest in that he worked with blood serum free from cells, and attempted to correlate the occurrences after the injection of animals with the action of the serum upon the cellular blood elements *in vitro*. Landois observed both the solution of hemoglobin and hemagglutination, and was led to regard the action of serum upon erythrocytes as the primary cause of death after transfusion. His conception of the mechanism is apparently twofold. On the one hand, he believed that when small quantities of blood were transfused, a formation of fibrin (stroma-fibrin) was initiated in the stroma of the injured erythrocytes which led to coagulation and thrombosis in the capillaries of the central nervous system and lungs. In the case of the transfusion of rabbit's blood into dogs he attributed death to embolism in the pulmonary vessels due to "Massenhafte Verklebung der Kaninchenzellen im Hundeblood"—or, in other words, to hemagglutination.

Ponfick and others have disputed the validity of Landois' conclusions, but the basic principles of his explanations have been upheld within recent years by workers who have gone over the same ground with the aid of more modern methods. Two careful re-

⁶⁸ A brief historical review of this work can be found in the paper of Coca, *Virchow's Arch. f. path. Anat.*, 1909, Vol. 196, p. 92.

searches have appeared during the last two years in which the problem has been approached by different routes, but in which the general conclusions show much agreement. Coca,⁶⁹ investigating the cause of death following the injection of washed blood cells into animals of different species, concludes that in these cases death is due to mechanical obstruction of the pulmonary circulation owing to agglutination of the injected cells. It is important to note, however, that he adds in his conclusions the following paragraph: "The mere presence of specific agglutinins does not suffice, in the injection of 'toxic' erythrocytes, to occlude the pulmonary circulation. The coöperation of another factor must be assumed—a factor probably found in the capillary walls."

Loeb, Strickler, and Tuttle⁷⁰ investigated the cause of death following the injection of normal dog and beef sera into rabbits. They correlated their animal experiments carefully with the action of the sera *in vitro* upon the blood elements of rabbits, and utilized the property of hirudin to inhibit the coagulation of blood, finding, in the case of dog serum, that injections of hirudin, while not always preventing death, at any rate prolonged life or necessitated an increase in the lethal dose. The conclusions of these authors are as follows: "Death following the injection of foreign serum is brought about by obstruction of the pulmonary circulation either by heaps of agglutinated erythrocytes or by fibrinous plugs. Dog serum and beef serum represent two different types. In the case of dog serum hemolysis of the blood cells of the recipient liberates substances attached to the stromata, which hasten coagulation. In consequence fibrin is formed which is carried into the pulmonary vessels and occludes them. In the case of beef serum death is due to hemagglutination."

The more recent understanding of the liberation of toxic bodies from blood cells by immune hemolytic sera, especially by the experiments of Friedemann cited above, have rendered it likely that a similar anaphylatoxin formation from the cells of the recipient may lie at the bottom of the toxic action of normal sera. And it is a fact, indeed, that such toxic sera are always hemolytic for the corpuscles of the susceptible animal.

An analysis of the toxic action of certain normal sera from this point of view has been made by Uhlenhuth and Haendel,⁷¹ who, in studying the necrotizing action of beef serum injected into guinea pigs, attribute this action of the serum to a "complex process depending upon the coöperation of complement," but not identical with the hemolytic mechanism. The toxic action of such serum, however, they separate from the necrotizing action, concluding that this

⁶⁹ Coca. *Virchow's Archiv*, Vol. 196, 1909.

⁷⁰ Loeb, Strickler, and Tuttle. *Virchow's Archiv*, Vol. 201, 1910.

⁷¹ Uhlenhuth and Haendel. *Zeitschr. f. Immunitätsforsch.*, Vol. 7, 1910.

is independent of complement, and more thermostable than either the mechanism causing necrosis or that responsible for hemolysis.

Recent studies of the writer⁷² on the toxic action of goat serum for rabbits have shown that, contrary to Loeb, Strickler, and Tuttle, hemagglutination and blood coagulation can be excluded as causes of death, and that, in agreement with Uhlenhuth and Haendel, the toxic action is due to a proteolytic action on the part of the serum not necessarily identical with the hemolysins, but producing from the protein of the recipient a poison similar to the anaphylatoxins. Unlike Uhlenhuth and Haendel, however, it seemed clear that the participation of alexin was definitely necessary—the process being probably entirely analogous to Friedemann's results with immune hemolytic (cytolytic) sera. The poisonous action of dissolved hemoglobin could be excluded. In principle, therefore, the toxic action of normal sera would seem to depend upon a mechanism similar to that of other anaphylactic phenomena.

Toxin hypersusceptibility, which is often acquired by animals in the course of immunization with diphtheria and tetanus toxin, is usually classified with anaphylaxis, indeed is often cited as the earliest observation of this phenomenon. However, it is by no means clear that the two conditions are actually analogous, since in the case of the toxins we are dealing with antigens which are not only toxic in themselves, but against which neutralizing antibodies are formed in the reacting animal. This last fact alone would separate toxin hypersusceptibility sharply from true protein-anaphylaxis in that entirely different reacting-mechanisms seem to be called into play by the two varieties of antigen. It will be necessary, therefore, to discuss toxin hypersusceptibility at some length.

Probably the earliest authentically recorded observation is that of von Behring,⁷³ who determined, both for diphtheria and tetanus toxins, that animals once inoculated with these poisons were occasionally more sensitive to them subsequently than were normal animals. He spoke of "Gift Ueberempfindlichkeit" as a property acquired by reason of a preceding injection, and the observation was further developed by Knorr⁷⁴ in 1895, and by v. Behring himself, in collaboration with Kitashima,⁷⁵ a few years later. These writers showed that guinea pigs which are treated repeatedly with small doses of diphtheria toxin may, under certain circumstances, not only fail to show immunity, but may even develop a susceptibility increased to such an extent that doses far too small to injure a normal animal will cause their death. Again, in the case of diphtheria toxin similar observations were made upon horses by both Salomonsen and

⁷² Zinsser. *Jour. Exp. Med.*, Vol. 14, 1911.

⁷³ Von Behring. *Deutsche med. Woch.*, 1893.

⁷⁴ Knorr. Quoted from Otto, "Dissertation." Marburg, 1895.

⁷⁵ Von Behring u. Kitashima. *Berl. klin. Woch.*, 1901.

Madsen⁷⁶ and by Kretz.⁷⁷ The last-named worker observed that horses that had been immunized with diphtheria toxin would often react to neutral mixtures of toxin and antitoxin by which normal horses were unaffected. This so-called "paradox phenomenon" was much discussed, and many theories advanced to explain it, a most ingenious adaptation of the side-chain theory being applied to it by Kretz⁷⁸ and by Wassermann.⁷⁹ They assumed that the partial immunization in such treated animals had in truth induced the formation of excessive receptors; that, in the stages of hypersusceptibility, however, these receptors had not yet been cast off from the cells. In consequence there was an excess of "sessile receptors"—by means of which the cell was rendered more exposed to toxin action than it was normally—it being still unprotected by the presence of freely circulating "antitoxin" receptors. The difficulties arising from the observation of similar hypersusceptibility in animals whose blood contained free antitoxin were disposed of by Wassermann by the convenient assumption of variations of affinity.

He assumed that the treatment with toxin, i. e., the intoxication, may induce a condition of higher affinity for the poison on the part of the sessile cell receptors, leading to a selective toxin-absorption by the cells and consequent greater susceptibility to injury. With Behring, he speaks of this as a "histogenic hypersusceptibility," implying an increased vulnerability of the tissue cells.

The analogy between these early observations and the phenomena which we now classify as anaphylaxis is unquestionably a striking one. However, it is doubtful, as Friedemann suggests, whether the two processes depend upon similar mechanisms. For, as we have seen in the case of the sensitiveness to toxin, we are dealing with primarily poisonous substances against which in the reacting animal neutralizing antibodies are found—a combination of conditions quite different from those with which we are confronted in hypersusceptibility against primarily harmless proteins. It is, of course, possible that the toxin hypersusceptibility is a true anaphylaxis against the toxin-protein—independent of the specifically poisonous nature of this substance. However, this is unlikely, since Löwi and Meyer⁸⁰ have shown that with tetanus toxin, the symptoms of such hypersusceptibility are not those of anaphylaxis, but of increased but characteristic tetanus poisoning. The fact that toxin hypersusceptibility cannot be passively transferred with the serum of a susceptible animal does not seem to us a good argument against its anaphylactic na-

⁷⁶ Salomonsen et Madsen. *Ann. de l'Inst. Past.*, 1897.

⁷⁷ Kretz. Quoted from Otto in "Kolle u. Wassermann Handbuch," *Ergänzungsband* 2, p. 232.

⁷⁸ Kretz. *Zeitschr. f. Heilkunde*, 1902.

⁷⁹ Wassermann. "Kolle u. Wassermann Handbuch," Vol. 4, p. 479.

⁸⁰ Löwi and Meyer. *Festschrift. Schmiedeberg Suppl., Arch. f. exp. Path. u. Therap.*, 1903, p. 355.

ture, since this, as we shall see, is equally impossible in the case of tuberculin susceptibility, which is in all probability a modified example of true anaphylaxis. Löwi and Meyer regard tetanus toxin hypersusceptibility as a "summation"—meaning thereby that it depends upon an alteration of the cells of the spinal cord because of traces of the poison retained in them. When the toxin was given intraneurally no antitoxin formation occurred, but the animals developed a marked hypersusceptibility in the course of several weeks, showing that here, unlike true anaphylaxis, specific antibodies play no part.

Not unlike toxin hypersusceptibility is that which is noticed in the case of certain medicinal substances. Such are the so-called idiosyncrasies against cocain, pilocarpin, morphin, quinin, and other drugs. These conditions have no direct relation to anaphylaxis, and, according to Hans Meyer,⁸¹ depend probably upon the chemical peculiarities of the tissues of the individual, such as calcium contents, etc. Hunt⁸² has also shown that poison susceptibility, in certain cases, may be influenced by the diet.

⁸¹ Meyer u. Gottlieb. "Experimentelle Pharmakologie," 2d Ed., Urban & Schwartzberg, pp. 520 *et seq.*, 1911.

⁸² Reid Hunt. *U. S. Pub. Health and M. H. S. Hyg. Lab. Bull.* 69, 1910.

CHAPTER XVII

BACTERIAL ANAPHYLAXIS AND ITS BEARING ON THE PROBLEMS OF INFECTIOUS DISEASE

IN the case of most serum reactions the original observations were made upon the sera of bacteria-immune animals, and later expanded into generalizations applicable to antigens as a class. This was the case with the phenomena of lysis, agglutination, and precipitation. In the case of anaphylaxis the reverse was true. The fundamental observations were made with non-bacterial antigens, but the thought that analogous observations could be made with bacterial proteins was an obvious one, and since the problem was one of altered susceptibility there was great promise that investigation of this subject might prove of profound significance for our knowledge of the pathology of infectious diseases.

Accordingly Rosenau and Anderson,¹ in one of their earliest researches, carried out experiments upon the sensitizing properties of bacterial proteins. They were successful in sensitizing guinea pigs with extracts of colon, tubercle, anthrax, and typhoid bacilli, with *Bacillus subtilis* extracts, and with those of yeast. In most cases they used considerable quantities of bacterial extracts and obtained but slight or moderate symptoms. However, their results were conclusive in showing that the anaphylactic experiment could be carried out with bacterial proteins and was, in every detail, analogous to the similar phenomena of serum anaphylaxis.

Not only could the basic experiment of active sensitization be carried out with these substances, but it was found that the reaction here, as in other cases, was specific, and that shock was followed by a period of "antianaphylaxis" or "immunity." Rosenau and Anderson suggested that the incubation time of many infectious diseases may be represented by the period necessary for the development of susceptibility after a first injection, and that the crisis of pneumonia might possibly find an explanation in the analogy with anaphylaxis.

The criteria governing the successful production of bacterial anaphylaxis were then studied especially by Kraus and Doerr,² Holo-

¹ Rosenau and Anderson. *U. S. Pub. Health and M. H. S. Hyg. Lab. Bull.* 36, 1907.

² Kraus and Doerr. *Wien. klin. Woch.*, No. 28, 1908.

but,³ Delanoë,⁴ and others, and the essential points of Rosenau and Anderson's experiments were confirmed. Although Kraus and Doerr succeeded in frequently sensitizing guinea pigs with a single injection of bacteria, this was not found to be the most favorable method for sensitization. Braun⁵ obtained entirely negative results by such a procedure, but this may well have been because in the first place single sensitization with bacteria is evidently irregular in result, and because Braun carried out his intravenous test-injection *slowly*, a technique by which Friedberger found later that shock could be avoided. Delanoë, in the main, confirmed the fact that bacterial sensitization was possible, but denied the specificity of the resulting anaphylaxis, in that he succeeded in producing shock in tubercle-sensitized guinea pigs with comparatively large amounts of typhoid, paratyphoid, and other bacilli, and conversely found typhoid-sensitized guinea pigs hypersusceptible to tubercle-injections. Other workers, however, notably Kraus and Doerr, Holobut, and Kraus and Admiradzibi,⁶ agree that the reaction is specific, at least in the same limits within which other serum reactions may be called specific.

Holobut then developed a technique of sensitization with bacteria more reliable than any which had been previously employed by other workers. He found that the most regularly successful results were obtained when he injected small quantities of bacteria (1/100 loopful) daily for ten days, subcutaneously, and tested with fairly large amounts (1-2 c. c. of an emulsion of the bacteria) intravenously about 3 weeks after the last sensitizing injection. This is in keeping with later experience, and in our own work with typhoid immunization in young goats we have found that anaphylactic reactions were not observed unless the goats had previously received several injections. A second injection never elicited symptoms.

It is not at all unlikely that this difference between serum sensitization and bacterial sensitization is due to the comparatively larger amounts of protein injected with very small volumes of serum than is the case with even the thickest bacterial emulsions. When larger sensitizing quantities of bacteria are used—which is often difficult because of the primarily toxic nature of some of the bacteria—a single sensitization gives positive results in guinea pigs more frequently than when the smaller amounts are used.

Since it was objected to many of the results at first obtained with bacterial sensitization that they might have been due to the primarily

³ Holobut. *Zeitschr. f. Immunitätsforsch.*, Vol. 3, 1909.

⁴ Delanoë. *C. R. de la Soc. de Biol.*, Vol. 66, 1909, pp. 207, 252, 348, 389.

⁵ Braun. Quoted by Bail and Weil, *Zeitschr. f. Immunitätsforsch.*, Vol. 4, 1910.

⁶ Kraus u. Admiradzibi. *Zeitschr. f. Immunitätsforsch.*, Vol. 4, 1910.

toxic nature of the bacteria or their extracts, it is important to note that Kraus and Doerr and later Kraus and Admiradzibi succeeded in well-controlled experiments in transferring bacterial anaphylaxis "passively" with the serum of previously sensitized animals—not only of the same, but of other species—(rabbit serum to guinea pigs). These experiments add the final link to the chain of complete analogy between bacterial and serum anaphylaxis.

This analogy was partly established and, in its completeness, clearly foreseen, when Friedemann's work upon the poisons produced from cells by hemolytic sera, and Friedberger's similar work upon serum precipitates, turned the trend of anaphylactic experimentation into new channels.

It will be remembered that, before this time, the toxic action of most bacteria (exclusive of "true toxin" producers like diphtheria and tetanus bacilli) had, since Pfeiffer, been attributed to the liberation of preformed "endotoxins" from the bacterial body during the process of lysis.

This idea is fundamental to the opinion of hypersusceptibility expressed by Wolff-Eisner⁷ as early as 1904.

The underlying concept of these ideas is really a morphological one in which the "endotoxin" is regarded as something present in the antigen which is set free by disintegration of the cell. In applying this to serum anaphylaxis Wolff-Eisner⁸ preserves this morphological simile in that he speaks of the dissolved protein antigen (serum, etc.) as "nur scheinbar gelöst" and "dass es erst durch die Lysine wirklich resorbierbar wird."

Indeed the sudden liberation of endotoxins by immune sera had been regarded by Pfeiffer and others as the cause of the rapid death often ensuing in *immunized* guinea pigs when more than a definite maximum of cholera spirilla or other organisms was injected. In all these opinions the basic conception was that certain bacteria contained a characteristic preformed poison (endotoxin) upon the pharmacological properties of which the peculiar symptoms caused by each organism depended.

The earliest unambiguous statements of a conception differing from this original view of the nature of bacterial endotoxins, and approaching the later conceptions of Friedberger, are found, we believe, in the work of Vaughan.⁹ In an article by him, published in 1908, Vaughan, after describing the incubation time occurring in man and animals after inoculation with typhoid bacilli, says: "The sickness begins when the animal body becomes sensitized and begins to split up the bacilli." By "splitting up" he means here, as in his

⁷ Wolff-Eisner. *Centralbl. f. Bakt.*, Vol. 37, 1904.

⁸ Wolff-Eisner. "Handbuch der Serum Therapie," p. 24, Lehmanns, München, 1910.

⁹ Vaughan. *Am. Jour. of Med. Sci.*, Sept., 1908.

other work¹⁰ on protein split products, not a mere liberation of pre-formed poisons, but a chemical (enzymotic) proteolysis by which a poisonous group of the bacterial protein-molecule is set free.

The essential difference of this point of view from the endotoxin theory at first sight seems a trivial one—in the one case liberation of a preformed poison molecule—in the other liberation of a poison by the breaking up of a molecule. The difference, however, is a fundamental one. For, in the earlier theory, the specific element of the toxemia was in the nature of the different poisons—whereas in the view of Vaughan the lysin which breaks up the protein molecule is alone the specific element, the formed poisons being concerned as non-specific and alike, whether produced from colon bacilli, tubercle bacilli, or egg white.

Friedberger,¹¹ finally, in 1910, repeating with bacteria his experiments upon “anaphylatoxin” liberation from specific precipitates, succeeded in obtaining such poisons in the test tube by allowing fresh guinea pig complement to act upon sensitized bacteria.

These results were confirmed by extensive experiments carried out soon after this by Friedberger¹² himself with a number of collaborators.

The results of these investigations may be summarized as follows:

1. The action of alexin upon sensitized or unsensitized bacteria yields toxic substances which, injected into normal guinea pigs, produce the characteristic symptoms of anaphylaxis, with frequent death and typical autopsy findings.

2. These poisons (“anaphylatoxins”) may be produced from any variety of bacteria, pathogenic and non-pathogenic.¹³ (The organisms used in the earlier experiments were *Vibrio metchnikovi*, the bacillus of tuberculosis, the typhoid, prodigiosus, and subtilis bacillus, and *Aspergillus fumigatus*.)

3. The successful production of the poisons depends intimately upon the relative amounts of antigen (bacteria) and alexin used, and upon the time and temperature conditions under which the exposures are made.

4. The poisons can be produced from boiled as well as from native bacteria.

Although unsuccessful with none of the bacteria with which experiments were carried out, different species yielded the poison with

¹⁰ Vaughan. *Zeitschr. f. Immunitätsforsch.*, Vol. 1, 1909.

¹¹ Friedberger. *Berl. klin. Woch.*, Nos. 32 and 42, 1910.

¹² Friedberger; Friedberger and Goldschmid; Friedberger and Szymanowski; Friedberger and Schütze; Friedberger and Nathan. *Zeitschr. f. Immunitätsforsch.*, Vol. 9, 1911.

¹³ Neufeld and Dold, comparing virulent and avirulent strains of pneumococcus in this regard, have found that the virulence of the race has no relation to its yield of anaphylatoxin. Indeed the anaphylatoxins from various bacteria seem to be qualitatively entirely alike.

varying degrees of intensity, though qualitatively the poisons were similar. *Bacillus prodigiosus*, though non-pathogenic, seems, in general, to be one of the most favorable micro-organisms for such experiments.

Since a clear understanding of Friedberger's basic experiments is essential to the further development of the theoretical conceptions which have been based upon them, it will be useful to insert here a protocol taken from his paper with Goldschmid.

Experiment VI. 30, VI, 1910. Ten 3-day agar cultures of typhoid bacilli washed up in salt solution—5 c. c. to 1/2 culture. Varying amounts of inactivated typhoid immune serum are added, the tubes brought to 11 c. c., 24 hours in refrigerator. 1, VII—Centrifugalized and to sediment added 4 c. c. guinea pig complement (active or inactivated), 24 hours in refrigerator. 2, VII—Centrifugalized and supernatant fluids injected into guinea pigs of 200 grams intravenously.

| Exp. No. | Amount of culture | Specific immune serum sensitiz. | Amount of complement | No. of animal | Symptoms | Result |
|----------|-------------------|---------------------------------|----------------------|---------------|---------------|--------------|
| 1 | 1/2 slant agar | 0 | 4 c. c. | G 61 | Severe | Dead 4 min. |
| 2 | 1/2 slant agar | 0 | 4 c. c. | G 64 | Slight | Dead 18 hrs. |
| 3 | 1/2 slant agar | 0 | 4 (heated 56° C.) | G 62 | No symptoms | Lives |
| 4 | 1/2 slant agar | 0 | 4 (heated) | G 63 | No symptoms | Lives |
| 5 | 1/2 slant agar | 1.0 | 4 | G 66 | Severe anaph. | Lives |
| 6 | 1/2 slant agar | 1.0 | 4 | G 69 | Very severe | Dead 4 min. |
| 7 | 1/2 slant agar | 1.0 | 4 (heated) | G 65 | 0 | Lives |
| 8 | 1/2 slant agar | 1.0 | 4 (heated) | G 68 | 0 | Lives |
| 9 | 1/2 slant agar | 0.1 | 4 | G 67 | Very severe | Dead 8 min. |
| 10 | 1/2 slant agar | 0.1 | 4 | G 70 | Very severe | Dead 11 min. |
| 11 | 1/2 slant agar | 0.1 | 4 (heated) | G 73 | 0 | Lives |
| 12 | 1/2 slant agar | 0.01 | 4 | G 71 | Very severe | Dead 2 min. |
| 13 | 1/2 slant agar | 0.01 | 4 (heated) | G 75 | 0 | Lives |
| 14 | 1/2 slant agar | 0.001 | 4 | G 72 | Very severe | Dead 5 min. |
| 15 | 1/2 slant agar | 0.001 | 4 (heated) | G 74 | 0 | Lives |
| 16 | | 1.0 | 0 | G 76 | 0 | Lives |
| 17 | | 0.1 | 0 | G 77 | 0 | Lives |
| 18 | 1/2 culture | 0 | 0 | G 79 | 0 | Lives |

From Friedberger and Goldschmid, *loc. cit.*, p. 402. (Changes made only in wording and omission of control 19.)

This series alone shows that, under the given conditions, 4 c. c. of alexin will produce the poison from 1/2 slant of typhoid bacilli, without sensitization (tubes 1 and 2), with sensitization ranging in degree from 1. c. c. to 0.001 c. c. of the given immune serum (tubes 5, 6, 9, 10, 12, and 14), and that inactivation of the alexin serum in all cases prevented the poison formation. Normal guinea pig serum alone, active or inactivated, the bacteria, or the immune serum alone were without toxicity in all of numerous controls.¹⁴

The experiments of Friedberger and his associates were rapidly

¹⁴ Injury of the animals by mere volume of injection can be definitely excluded. The writer has frequently injected 5 to 6 c. c. of salt solution into guinea pigs of 200 to 300 grams without symptoms in any way resembling anaphylaxis.

confirmed by Neufeld and Dold,¹⁵ Kraus,¹⁶ Ritz and Sachs,¹⁷ and many others,¹⁸ and, though the conditions under which the anaphylatoxin formation took place were defined with slight variation by different workers, the essential features of Friedberger's claims were upheld.

As was to be expected, it was soon found that instead of the prolonged exposures at refrigerator temperature the poisons could be obtained more rapidly by digestion for shorter periods in water¹⁹ baths at 37° C. And with this method accurate studies on the relations between time of exposure and proportions of reagents (antigen, sensitizer, alexin) were made, relations the importance of which was apparent from Friedberger's first studies. The outcome of this work was as follows: 1. There are a definite minimum and a definite maximum quantity of bacteria from which anaphylatoxin can be produced by a given fixed quantity of guinea pig serum. Thus, in one of the experiments of Friedberger and Goldschmid, 4 loopsful of typhoid bacilli with 4 c. c. of complement produced a fatal poison, 24 loopsful with the same amount produced none. (In some of the writer's²⁰ experiments with typhoid bacilli a similar principle of proportions was evident, though much larger quantities of typhoid bacilli could be successfully used if the time of exposure at 37° C. was prolonged.) 2. If sensitized bacteria are used an excess of sensitization, beyond a definite limit, weakens the formation of anaphylatoxin. It may be permitted to illustrate this with a protocol of one of the writer's experiments with typhoid bacilli, since, though merely confirming the principle laid down by Friedberger, it included a careful titration of the bactericidal contents of the anti-typhoid serum.

TITRATION EXPERIMENT WITH TYPHOID-IMMUNE SERUM

Rabbit 79

| Dilution of serum | Agglutination | Bactericidal titre with modified Stern-Korte method |
|-------------------|---------------|---|
| 1:100 | +++ | 480 colonies |
| 1:200 | +++ | 556 colonies |
| 1:500 | +++ | 750 colonies |
| 1:1,000 | ++ | Over 10,000 colonies |
| 1:2,000 | ± | +++++ |
| 1:5,000 | — | +++++ |
| 1:10,000 | — | +++++ |

¹⁵ Neufeld and Dold. *Berl. klin. Woch.*, No. 2, 24, 1911; *Arb. a. d. kais. Gesundheits amt.*, Vol. 38, 1911.

¹⁶ Kraus. *Zeitschr. f. Immunitätsforsch.*, Vol. 8, 1911.

¹⁷ Ritz u. Sachs. *Berl. klin. Woch.*, No. 22, 1911.

¹⁸ Izar. *Zeitschr. f. Immunitätsforsch.*, Vol. 11, 1911.

¹⁹ Friedberger u. Mita. *Zeitschr. f. Immunitätsforsch.*, Vol. 10, 1911.
See also Dold, "Das Bakterien Anaphylatoxin," Fischer, Jena, 1912.

²⁰ Zinsser. *Jour. Exp. Med.*, Vol. 17, 1913.

Two-tenths c. c. of this serum added to 1 c. c. of typhoid filtrate gave a very slight clouding in about 15 minutes.

ANAPHYLATOXIN EXPERIMENTS

| Number in series | Typhoid bacilli | Amount of inactive serum antityphoid | Amount of complement | Weight of animal | Result |
|------------------|----------------------|--------------------------------------|----------------------|------------------|----------------------|
| 1 | $\frac{1}{10}$ slant | 5.0 c. c. | 4 c. c. | 215 gm. | Very sick, recovers |
| 2 | $\frac{1}{10}$ slant | 3.5 c. c. | 4 c. c. | 200 gm. | Typical death 2 min. |
| 3 | $\frac{1}{10}$ slant | 3.0 c. c. | 4 c. c. | 198 gm. | Typical death 2 min. |
| 4 | $\frac{1}{10}$ slant | 2.0 c. c. | 4 c. c. | 225 gm. | Typical death 2 min. |
| 5 | $\frac{1}{10}$ slant | 1.0 c. c. | 4 c. c. | 200 gm. | Sick, recovers. |

The weak character of the antiserum used, and the fact that, in this experiment, the digestion was at 0° to 5° C., explain the failure to obtain a strong anaphylatoxin with 1 c. c. of sensitizing serum.

The negative experiment resulting from a too vigorous sensitization is practically a corollary of the next point ascertained by Friedberger, namely, that:

3. With constant amounts of reagents a too prolonged exposure at 37° C. will result in failure to obtain the poison.

How are we to explain these experimental results? The first of the three—namely, the fact that an excess of bacteria inhibits the formation of anaphylatoxins—seems to the writer most easily explained by accepting the views of Bordet on the manner of the union of an antigen with its antibody. For, unlike the opinion of Ehrlich, who assumes a union of the two according to the laws of multiple proportions, Bordet²¹ believes that the distribution of serum substances upon an antigen is such that the entire amount of antibody is distributed equally among the antigenic elements. In the case of an excess of bacteria, as in these experiments, therefore, the quantity falling to each unit is insufficient, at least in the time of exposure here practiced, to accomplish the cleavage necessary for poison production.

As regards the second and third point—the failure of producing anaphylatoxins if, on the one hand, too intense sensitization was employed—or, on the other, the time of exposure was too prolonged—these seem to indicate that anaphylatoxin is not the end product of the complement action, but rather an unstable intermediate substance which, once formed, is rapidly further decomposed (“abgebaut”) into non-toxic derivatives.

²¹ Bordet. *Ann. de l'Inst. Past.*, 17, p. 161, 1903.

Indeed, Neufeld and Dold,²² in experiments with the cholera spirillum, found that whenever lysis was permitted to proceed as far as the actual disintegration and granulation of the bacteria no poisonous substances were obtained. They conclude from this that rapid lysis actually prevents the production of the poison, and that the anaphylactic antibody has no relation to the bacteriolytic sensitizer. They fortify this opinion by experiments in which they easily obtained powerful poisons with pneumococci, organisms which are but slightly, if at all, subject to actual lysis. They suggest identity of the anaphylactic antibody with the opsonins, or possibly with the "Bordetsche Antikörper" of Neufeld. This latter conclusion does not seem valid, since the mere fact that one micro-organism undergoes lysis and another does not is not necessarily an argument for a difference in the sensitizers produced in animals by immunization with these bacteria. It may, and probably does, depend upon variations in the ease of disintegration of the different cell-bodies, and, as a matter of fact, not many bacteria undergo actual complete lysis as easily as does the cholera spirillum. Moreover, there is much evidence in favor of the so-called "unitarian" point of view, which holds that no fundamental structural and functional differences between the various heat-stable antibodies—sensitizers (amboceptors), precipitins, immune opsonins (bacteriotropins), and the so-called "Bordet" alexin-fixing antibodies—have as yet been proved.

However this may be, it seems conclusively established that a too vigorous and prolonged action of the antibody-alexin complex upon the bacterial protein does not yield poisons—and that, since less vigorous sensitization or early interruption of the exposure will lead to positive results, the mechanism is one of rapid poison formation with equally rapid further decomposition into a non-toxic substance. In some cases this is more rapid than in others. In Neufeld and Dold's experiments with cholera spirilla the exposure of 2 loopsful of the organisms sensitized with 0.02 antiserum and treated with 2 c. c. of alexin resulted in complete lysis and failure of demonstrable anaphylatoxin in 2 hours at 37° C. In some of the writer's experiments with typhoid bacilli the most regular positive results were obtained when the exposures at 37° C. were prolonged to several hours and powerful poisons were determined even after as long as 15 hours at 37° C. An example of such an experiment is given below, since we believe that in the apparent stability of the typhoid anaphylatoxins and the wide range of quantitative relations within which the poison was successfully obtained, it forms a strong argument in favor of Friedberger's theory of the rôle played by these poisons in diseases like typhoid fever.

²² Neufeld and Dold. *Loc. cit.*

EXPERIMENT II ²³

The materials used were *Bacillus typhosus* 65, typhoid-immune rabbit serum (from rabbit A),²⁴ inactivated at 56° C., and fresh guinea-pig serum as complement. The injected guinea pigs weighed from 150 to 225 grams.

| No. in series | Amount of bacteria | Sensitized with inactive serum 1 hr. at 37.5° C. | Time of exposure to complement at 37.5° C. | Amount injected | Result |
|---------------|--------------------|--|--|-----------------|---------------------------|
| 1 | 1½ slant | 1 c. c. | 15 hrs. | 4 c. c. | Very sick, recovers |
| 2 | 1 slant | 1 c. c. | 15 hrs. | 4 c. c. | Typical death in 8½ mins. |
| 3 | 2 slants | 1 c. c. | 15 hrs. | 4 c. c. | Typical death in 5 mins. |
| 4 | 2 slants | Not sensitized | 15 hrs. | 4 c. c. | Typical death in 5½ mins. |
| 5 | 3 slants | 1 c. c. | 15 hrs. | 2.5 c. c. | Typical death in 4 mins. |
| 6 | 3 slants | 4 c. c. | 15 hrs. | 4 c. c. | Typical death in 4 mins. |
| 7 | 3 slants | Not sensitized | 15 hrs. | 4 c. c. | Typical death in 2 mins. |
| 8 | 8 slants | 1 c. c. | 15 hrs. | 4 c. c. | Typical death in 5 mins. |
| 9 | 8 slants | Not sensitized | 15 hrs. | 4 c. c. | Typical death in 4 mins. |
| 10 | 12 slants | 1 c. c. | 15 hrs. | 4 c. c. | Very sick, lives |
| 11 | 12 slants | Not sensitized | 15 hrs. | 4 c. c. | Slightly sick |

Similar in significance to the points just considered also is the experiment of Friedberger and Szymanowski with *Vibrio metchnikovi* (confirmed by the writer with typhoid bacilli) that, although sensitized and unsensitized bacteria will yield anaphylatoxin with almost equal intensity, the poisons are produced from the sensitized bacteria with far greater speed than from the latter. The difference between the two, in fact, is probably one of degree only, since in experiments without the addition of specific antiserum the bacteria are nevertheless slightly sensitized by the normal antibody present in the guinea-pig serum.

That the production of the poison can under no circumstances be regarded merely as a giving up from the bacterial cell of preformed endotoxins under the influence of lytic substances which produce greater permeability of the cell membrane was shown by Neufeld and Dold, who extracted bacteria with lecithin salt solution and pure salt solution, and from these extracts (but moderately toxic in themselves) produced typical anaphylatoxins by the action of complement. The matrix of the poison thus is shown by direct experiment to be a soluble ingredient of the bacterial cell.

It was further shown by Friedberger and Nathan that the conditions prevailing in the test tube experiment in truth represent the processes taking place within the animal body. This they accomplished by injected bacterial emulsions into the peritoneal cavities of

²³ Zinsser. *Loc. cit.*

²⁴ This serum had an agglutinating titre of 1:8,000 for *Bacillus typhosus* 65.

guinea pigs, killing the animals after several hours and examining the peritoneal exudates for their toxic properties. Centrifugalized, cleared of bacteria, and injected intravenously into other guinea pigs, these exudates produced the typical acute symptoms characteristic of the poisons obtained in test-tube experiments.

It was on these premises, then, that Friedberger²⁵ was led to formulate his views of the nature of bacterial infections, which give promise of introducing a new understanding of these diseases. It has been shown in the researches upon serum anaphylaxis that the injection of small quantities of a foreign protein may produce reactions of temperature which simulate very closely those prevailing in infectious diseases, and variations in the quantities injected, the path of administration, and the interval between injections may lead to conditions, local and systemic, which may affect, more or less profoundly, many different organs and tissues of the body. These matters we have considered in the general discussion of anaphylactic phenomena. Friedberger now suggests that we may regard bacterial infection, after all, as the presence in the body of a living foreign protein—in this case varying in distribution and quantity by reason of the particular invasive properties of the given germ and the balance between these and the resistance of the host. It is not necessary, therefore, to assume that the character of the disease is determined by the existence of different preformed “endotoxins.” He believes that we may justly assume that the toxic substances appear only after proteid cleavage of the bacterial bodies has been initiated by the action upon them of the serum components, and that the apparent specificity of the poisons, or differences between the toxemic manifestations of various diseases, may depend, not on differences in the pharmacological actions of these poisons, but rather upon variations in the invasive properties of the bacteria, both as concerns their quantitative distribution and their accumulation and localization in the infected body.

If we leave out of consideration bacteria which, like the diphtheria bacillus, produce true secretory poisons, it would be the ability to gain a foothold in the body, the degree of invasive power, the predilection in the choice of a path of entrance, and the specific local accumulation upon which the speed and quantity of anaphylatoxin production and absorption would depend, and which consequently would give character to variations in the clinical pictures of different diseases. Besides simplifying considerably our comprehension of bacterial toxemia this point of view again brings out the great importance of the work of Vaughan, and of Vaughan and Wheeler, on the non-specific poisonous fraction obtained by hydrolysis of bacterial and other proteids.

²⁵ Friedberger. *Loc. cit.*; also *Deutsche med. Woch.*, No. 11, 1911; *Berl. klin. Woch.*, No. 42, 1911.

To support this assumption Friedberger points out the similarity in the clinical manifestations of several diseases in which the inciting bacteria are biologically very different, but in which the distribution and invasive properties are alike. For instance, lobar pneumonia caused by the pneumococcus is clinically very similar to that caused by the Friedlander bacillus, though the micro-organisms inciting them are extremely unlike each other. He draws a similar parallel between true cholera and cholera nostras, and we may add another striking example in the great similarity existing clinically between the various forms of acute and subacute septicemia in which a definite bacteriological diagnosis can rarely be made except by blood culture.

Conversely the same micro-organism may call forth diseases which clinically apart from the purely local manifestations are very dissimilar, according to the localization and distribution of the bacteria.

Granted that we accept this view, then the subsidence of the disease might depend merely upon limitation of the supply of antigen, as the increasing bactericidal action of the blood constituents comes into play, and upon the consequent diminution of the anaphylatoxin. For, as the bacteria diminish and the sensitizer increases, a changed proportion between them is established which, finally, as experiment has shown, results in a failure of anaphylatoxin production. For, although experiments have shown that, within a wide latitude of relative proportions of bacteria and antibody, anaphylatoxin can be formed, beyond this range an excess of one or the other element eventually will prevent their formation.

Infectious disease, then, according to this point of view, represents merely the reaction of the body against a foreign protein, the bacteria. These gain a foothold in the body, and at first, during the so-called incubation time, cause no symptoms, since the slight amount of bacterial destruction with correspondingly slight cleavage of the bacterial protoplasm liberates too small an amount of anaphylatoxin to incite noticeable deviations from the normal condition. As these slight quantities of bacterial cleavage products are absorbed, however, a reactionary formation of specific antibody occurs. Meanwhile, also, the foreign protein increases and is distributed by bacterial growth. In consequence of these parallel processes changes of proportion between the reacting substances are created and a constantly greater amount of anaphylatoxin is liberated and the disease progresses. This may kill the patient if the proportions become such that the amount of poison formed exceeds the lethal dose. At any rate, the symptoms may vary and fluctuate according to the relations maintained between the reacting bodies, modified somewhat by the supply of alexin or complement. If recovery is to take place the amount of antibody (sensitizer, amboceptor) may become so great

that the bacteria are subjected to rapid destruction, the chemical cleavage of their bodies taking place so vigorously that practically no anaphylatoxin is distributed and vigorous phagocytosis is initiated. Finally the antigen is completely removed. On the other hand, an excessive increase of the bacteria or a defective supply of alexin might also lead to a final cessation of the formation of anaphylatoxin; in this case, however, we would expect death by the metabolic disturbance occasioned by the life processes of the great masses of bacteria. It is not unthinkable, moreover, that the bacterial enzymes in such a case might produce substances comparable to the anaphylatoxins from the destroyed tissues of the host.

It is perfectly true, as Friedberger says, that on the basis of this theory, rendered so likely by experimental fact, the assumption of the existence of endotoxins to explain the various manifestations of infectious disease is not necessary. The poisons, according to the view just outlined, are alike and non-specific. It is the reaction bodies, the sensitizers, induced by the bacterial protein which in each case are these specific elements.

While it is not necessary to assume specific endotoxins, however, it is not possible on present evidence to entirely exclude the participation of such substances in the genesis of infectious disease. The rapid toxic action of bacterial extracts obtained in various ways has been taken to argue in favor of this.

It is a difficult question to settle, and must undoubtedly remain an open one until a method is found by which crucial experiments can be formulated. Since Neufeld and Dold have succeeded in producing anaphylatoxin from bacterial extracts, the primary toxic action of every bacterial extract, however rapidly produced from the bacteria, can be regarded as possibly furnishing merely an antigen for anaphylatoxin production, and indeed such a supposition is rendered more likely by the almost invariable incubation time following upon the administration of endotoxic extracts, even when they are introduced directly into the circulation. Pfeiffer²⁶ himself still believes in specific endotoxins, basing his opinion on the individually characteristic nature of the infections caused by supposedly endotoxic bacteria. The differences in the degrees of toxicity, moreover, of extracts obtained by the same technique from different microorganisms would certainly tend to add some weight to his argument. We need only to recall to memory the greater toxicity of bouillon culture extracts of *B. dysenteriae* Shiga-Kruse as compared with similar extracts of *B. dysenteriae* Flexner or Hiss-Russell, or the similar difference between typhoid and colon extracts. Altogether the problem is an involved one, for the recent claims of Kraus,²⁷

²⁶ R. Pfeiffer. "Über Bakterien Endotoxine, etc.," Weichhardt's *Jahresbericht*, Vol. 6, p. 29, 1910.

²⁷ Kraus. *Monatschr. f. Gesundheitspflege*, No. 11, 1904.

Doerr,^{28 29} and others of having discovered true (antitoxin-forming) soluble toxins³⁰ in such cultures as those of cholera, dysentery Shiga, and typhoid bacilli add another complication. The present status of the question, it seems to us, may be summed up as follows: It may probably be accepted as a fact that anaphylatoxin production occurs and accounts for toxemia, altogether or in part, in all diseases in which bacteria invade the tissues or circulation; in addition to this, soluble toxins produced by the bacteria still living and uninjured may add a further specific element to the condition—in some diseases; whether or not specific preformed endotoxins participate in the production of bacterial toxemia cannot be definitely stated. It is not, however, a necessary assumption.

It still remains for us to consider certain experimental facts which have had some influence upon extending and altering the conceptions of anaphylatoxin formation which we have just outlined. In the earlier work of Friedberger, Neufeld and Dold, and others the poisons were formed from the bacteria by the action of alexin at low temperatures. This suggested the possibility that the alexin fractions—"Endstück" and "Mittelstück"—might not both be involved in the reaction, since, from previous studies, it was known that at low temperatures the midpiece (the globulin fraction) was bound, but that the end piece did not become active until the temperature was increased. This point was, therefore, made the object of a special investigation by Friedberger and Ito,³¹ who found that neither fraction alone would suffice, but that bacterial anaphylatoxins were formed only under the influence of the intact whole alexin, or by that of the two fractions, reunited after separation.

Because of the reasoning along which the investigations of anaphylatoxin formation were developed, it is not surprising that it seemed self-evident that the matrix of the poison was represented by the bacterial protein—the antigen of the lytic complex. The only fact which, in the earlier experiments, might have cast some doubt upon this was the ease with which anaphylatoxins were produced from boiled bacteria and precipitates and from such very insoluble organisms as the tubercle bacillus.

Such vague suspicion becomes a very definite doubt, however, in the light of the experiments of Keysser and Wassermann.³² Keysser and Wassermann utilized the fact that certain serum elements may be absorbed out of serum if this is shaken up with such indifferent suspensions as barium sulphate or kaolin (aluminium

²⁸ Kraus u. Doerr. *Wien. klin. Woch.*, No. 42, 1905.

²⁹ Kraus. "Kraus u. Levaditi Handbuch," Vol. 1, p. 180.

³⁰ Exotoxins.

³¹ Friedberger and Ito. *Zeitschr. f. Immunitätsforsch.*, Vol. 11, 1911.

³² Keysser and Wassermann. *Folia Serologica*, Vol. 7, 1911; *Zeitschr. f. Hyg.*, Vol. 68, 1911.

orthosilicate).³³ They therefore substituted these insoluble substances for antigen, allowed them to absorb serum constituents, assumed by them to be amboceptor, out of normal and inactivated immune sera, and then allowed complement or alexin to act upon the "sensitized" kaolin.

In this way they obtained active and powerful anaphylatoxin, and claim, in consequence, that the matrix of the poison is not in the bacterial antigen, but in the sensitizer or amboceptor, which is mechanically absorbed by the bacteria (as by the kaolin), and thus made amenable to the alexin action.

The experiments of Keysser and Wassermann have found confirmation in the hands of other investigators, although the results of Neufeld and Dold, as well as our own, with this method were far more irregular than were those of Keysser and Wassermann. Neufeld and Dold³⁴ and Friedberger³⁵ suggest that the horse serum absorbed by the kaolin may act as an antigen itself, and is acted upon by normal sensitizer present in the guinea pig serum. This is in keeping with the well-known fact that small amounts of sensitizers to many varieties of foreign proteins are present in normal serum, and is further borne out by the fact that Neufeld and Dold, unlike Keysser and Wassermann, were never able to produce anaphylatoxin by allowing the alexin alone to act upon kaolin—without previous absorption of horse serum.

We say "never," though the protocols of Neufeld and Dold³⁶ show a single successful experiment. This they explain, however, by assuming the accidental presence of some antigen in the alexic serum. That is, the entire complex, antigen, sensitizer, and alexin, is assumed to have been present in this particular guinea pig serum. The same explanation may be applied to the occasional inherent toxicity which develops in normal guinea pig sera on standing. Whether the above complicated explanation is necessary or whether we may assume an autolytic process in the guinea pig serum by which anaphylatoxin-like substances are formed is an open question.

At any rate, it has been shown that, even with bacteria, the action of alexin is not the only way in which acute poisons may be obtained from them. And, indeed, if we look upon the action of alexin as analogous to that of an enzyme—an assumption for which we have much supporting evidence, we may well expect that other methods of proteolysis will give similar toxic cleavage products. And various methods of bacterial autolysis have ac-

³³ Kaolin emulsions will absorb amboceptor only out of diluted serum. Out of concentrated serum complement is completely absorbed. Friedberger u. Salecker, *Zeitschr. f. Immunitätsforsch.*, Vol. 11, 1911; Zinsser, from *Journ. Exp. Med.*, Vol. 18, 1913.

³⁴ Neufeld and Dold. *Loc. cit.*

³⁵ Friedberger and Salecker. *Zeitschr. f. Imm.*, Vol. 11, 1911.

³⁶ Dold. *Loc. cit.*

tually yielded such results. Thus Neufeld and Dold obtained poisons by digesting typhoid bacilli, cholera spirilla, and other micro-organisms for several hours in salt solution, lecithin salt solution, and inactivated guinea pig sera. Their extracts killed guinea pigs within several hours. Rosenow³⁷ has even succeeded in obtaining acutely toxic substances which caused typical anaphylactic death in guinea pigs by suspending pneumococci, typhoid bacilli, and other bacteria in salt solution at 37° C. for varying periods, and the writer,³⁸ though never producing acute death, was able to cause typical anaphylactic shock in isolated cases with similar salt solution extracts of typhoid bacilli. It is not impossible that poisons obtained in this way are formed by autolysis due to proteolytic enzymes of the bacterial cell.

In cases in which bacteria, suspended in salt solution and other indifferent fluids, represent the only source of protein present it must, of course, be assumed that they are the substratum or matrix of the anaphylactic poison. They are also, of course, to be regarded as the source of the poison in such experiments as those of Vaughan, in which the poison was produced by chemical hydrolysis of the bacterial bodies. In the case of anaphylatoxin production by fresh serum in the presence of bacteria, kaolin, precipitates, etc., the question is much more complex.

As we have stated before, it is only natural, considering our previous knowledge of bacteriolysis in serum, that the first conclusion arrived at should look for the source of the poisons in the bacterial cells. The doubt which has been cast upon this assumption by the work of Keysser and Wassermann and others, however, rests upon a sufficiently sound experimental basis to prevent our absolute acceptance of this view. Jobling and Peterson³⁹ have recently carried out experiments which may serve to throw much light upon anaphylatoxin. They believe that, by the ordinary technique of anaphylatoxin production with bacteria and serum, most of the toxic substances originate from the serum proteins. The bacteria act merely by removing the anti-ferments from the serum, thereby setting free the ferments normally present in the serum, and permitting them to act upon the serum proteins. The result is cleavage and the production of toxic split products. This would explain such results as those of Keysser and Wassermann. Jobling and Peterson have supported their contention by experiments in which they have obtained typical anaphylatoxins by removing serum anti-ferments with chloroform, kaolin, and agar. They have further shown that emulsions of bacteria actually do remove anti-ferments from fresh serum, and that

³⁷ Rosenow. *Jour. Inf. Dis.*, Vol. 9, 1911; Vol. 10, 1912.

³⁸ Zinsser. *Loc. cit.*

³⁹ Jobling and Peterson. *Jour. of Exp. Med.*, June, 1914.

the bacteria used in the process become more resistant to tryptic digestion in consequence.

This does not necessarily weaken the force of Friedberger's view of infectious disease. For, whatever the source of the toxic substances, the result is still the same. Wherever proteolysis takes place, and certain quantitative relations between cleavage, energy, and substratum exist, it seems toxic bodies may be liberated.

And the result of such proteolysis, at some stage of the process, yields apparently the same non-specific toxic substance, whatever the particular nature of the proteolysis and whatever the variety of the original protein matrix.

CHAPTER XVIII

THE CLINICAL SIGNIFICANCE OF ANAPHYLAXIS

SERUM SICKNESS

WE have mentioned that Rosenau and Anderson attacked the problem of hypersusceptibility primarily in the hope of casting light upon the nature and cause of the distressing symptoms which in human beings often ensue upon the injection of diphtheria antitoxin. It has been one of the staple objections of lay opponents to the use of antitoxins that the injections are apt to cause severe illness and occasionally death, and indeed a few cases are on record in which sudden death has followed the first injection of diphtheria antitoxin. Since it was known by accumulated clinical experience as well as by experiments like those of Bertin,¹ of Johannesen,² and others that the harmful effects were not dependent upon the antitoxin contents, but could be produced by injections of normal horse serum, it was but natural to bring these ill effects into analogy with the phenomena of hypersusceptibility. A large number of references to such antitoxin illness or SERUM SICKNESS have appeared in the literature since the first beginnings of the therapeutic use of sera, yet no careful analysis of the condition was made until von Pirquet and Schick,³ in 1905, published their studies.

As a rule the results of serum injection have been mild and without danger, though sufficiently frequent and troublesome to call for thorough study and attempts to discover the prophylactic measures. As stated above, a few cases are on record in which sudden death has followed a single first injection. There are no reports in the literature known to us, however, of fatalities after second injections, although not infrequently such cases have taken on alarmingly serious aspects.

The percentage of incidence and the variety of symptoms have been the subjects of many reports. The most frequent and striking single occurrence has been an urticarial rash. Rolleston,⁴ in a large

¹ Bertin. *Gaz. Méd. de Nantes*, 1895. Quoted from Levaditi.

² Johannesen. *Deutsche med. Woch.*, No. 51, 1895.

³ Von Pirquet u. Schick. "Die Serum Krankheit," Deuticke, Leipzig, 1905. Also *Münch. med. Woch.*, 53, p. 67, 1906.

⁴ Rolleston. *The Practitioner*, Vol. 74, 1905.

series of cases, found urticaria in all but 17 of 289 cases of serum eruptions occurring between the first and tenth days after injection, and in all but ten of ninety-four later eruptions.

Rashes occurred in from 69.4 to 81.9 per cent. of the 600 anti-toxin cases which Rolleston reports.

Joint pains commonly accompany the appearance of the rash, and frequently there is adenitis, involving the glands adjacent to the point of injection, and even remotely in the submaxillary, axillary, or inguinal glands. Albuminuria is quite common, and with it oliguria and relative concentration. Fever is rarely absent, though usually slight, together with general malaise. Rolleston in his purely clinical study does not classify his cases into those reacting after a first injection and those showing symptoms after repeated treatment. He states, however, that the serum-reaction may be extremely severe in cases of "relapse or second attack of diphtheria" in which urticaria with "pronounced edema surrounding the wheals, vomiting, rigors, and collapse may ensue within a few hours of injection," and further asserts that these severe symptoms are more apt to follow upon large than upon small doses.

Von Pirquet and Schick have studied the condition with careful reference to a comparison between the symptoms occurring in subjects after a first injection of serum and those following upon repeated treatments. Their studies revealed the very important fact that the ill effects following a second injection were not only more severe than those occurring after the first injection, but developed after much shorter periods of incubation. In the ordinary "first injection" case the symptoms appear usually in from one to twelve days. After a second injection this incubation period may be considerably shortened and symptoms may appear in from five to seven days, the local and general reactions being much more marked than those subsequent to a first injection. Indeed, in some of the cases reported they may attain very alarming degrees of severity. This is the so-called accelerated ("beschleunigte") reaction of von Pirquet and Schick, and is different from the "first injection" symptoms only in its greater severity and speedier onset. In addition to this, however, the "second injection" cases may show a train of immediate symptoms⁵ (sofortige Reaktion), which occur within twenty-four hours after injection, and are characterized by marked local erythema and edema with often urticaria and constitutional disturbance. Both reactions may occur in the same individual, the "accelerated" reaction setting in as the "immediate" reaction subsides.

Again, one reaction or the other may occur alone. The analogy between the immediate reaction and the anaphylaxis of animal experiment is obvious. The cases may be classified on the basis of

⁵ Rankin in the *Lancet*, Dec., 1911, reports a case of "immediate" reaction 15 minutes after injection.

these reactions, according to von Pirquet and Schick, the nature of the reaction being, within certain limits, determined by the interval ensuing between the first and the second injection. Thus, when the interval was twenty-one days or less the immediate reaction alone was noticed. When the interval was between two and six months both the immediate and accelerated reactions were present, and when the interval was still longer (seven months or more) the accelerated reaction alone was present. Isolated exceptions to this are noted in the series of sixty-one cases so reported.

Currie,^{6 7} who has made similar studies, confirms the results of von Pirquet and Schick in all essentials, and agrees with their statement that the nature of the reaction is chiefly dependent upon the interval between injections.

That the entire train of symptoms, as well as the mere fact of their dependence upon an injection of a foreign protein, rather than upon the antitoxin itself, force upon us the analogy with anaphylaxis is clear. Moreover, this analogy becomes almost an identity when we can show, as von Pirquet and Schick have done, that the first injection has apparently sensitized the subject, in that the second administrations are fraught with more violent and serious reactions, dependent to a great extent, as in experimental anaphylaxis, upon the time intervening. If serum-sickness is truly an anaphylactic phenomenon, however, it is still by no means clear why symptoms should at all ensue after the first injection. Many explanations have been offered for this; none of them, however, from the very nature of the problem itself, can be finally accepted as proved. Two possible explanations appear from the experimental work of Rosenau and Anderson quoted above. These workers, we have seen, showed among other things that the state of hypersusceptibility could be transmitted from mother to offspring, and that sensitization by way of the intestinal canal was at least possible. Both of these factors may have determinative significance in the present case.⁸ There may be, because of such conditions, a pre-existent sensitization which, especially in cases of accidental injection of the antitoxin directly into a small vein (an accident probably not infrequent in deep muscular injections), may possibly explain the few instances of sudden death following the first antitoxin injection and the isolated instances of "immediate" reaction following "first" injections. Rosenau has also suggested recently that sensitization may be unconsciously acquired against various forms of protein by absorption through the lungs of the organic matter carried in the expired breath of animals. In this way possibly hyper-

⁶ Currie. *Jour. of Hyg.*, Vol. 7, 1907.

⁷ See also Goodall, *Jour. of Hyg.*, 7, 1907.

⁸ Regarding intestinal sensitization see also Richet, *C. R. de la Soc. de Biol.*, Vol. 70, 1911; Lesne et Dreyfus, *C. R. de la Soc. de Biol.*, Vol. 70, 1911.

susceptibility against horse protein may be acquired and subsequently be expressed by a reaction to the first injection of antitoxin.⁹ While this must be considered a possibility, however, not all investigators are ready to accept it and its significance is at present very uncertain.

In the ordinary case of serum-sickness after first injection, however, the long incubation time elapsing between the injection of the serum and the onset of symptoms, often more than 10 days, would, it seems to us, tend to argue against a previous hypersusceptible state of the patient. On the other hand, we have learned since¹⁰ the earlier studies of Eisenberg, von Dungern, and others that foreign proteins injected into rabbits may be excreted very slowly, and that even after the formation of antibodies (precipitins) the antigen may still be demonstrable in the blood serum of the rabbit. Thus at periods eight to twelve days after the injection of comparatively large amounts there is often present in the same individual both antigen and its specific antibody side by side, and the essential conditions for the production of anaphylatoxin are thus established. That the two bodies do not, as a rule, unite in such serum in quantities sufficient to be demonstrated by alexin fixation has been discussed in another place, but this by no means excludes a gradual, slow union of small amounts of antigen with antibody, consequent fixation of alexin, and the liberation of anaphylatoxic products. In fact, although there does not seem at present to be any way to bring experimental proof to support it, it seems very likely that a slow splitting of the antigen begins by virtue of the normal antibody, and as, in the course of eight to ten days, the antibody appears in relatively larger amounts, the toxic products of the reaction are sufficient to give rise to symptoms. Such a point of view is supported only by the experimental knowledge that antibody may appear in considerable concentration before the antigen has disappeared from the circulation, and upon the facts we know concerning the toxic substances which arise from the union of two such reagents subjected to the influence of alexin.

In fact, it seems likely that this process of antibody formation may represent merely an emergency mechanism for the purpose of ridding the body of foreign dissolved proteins which have penetrated into the circulation, cannot diffuse unchanged through the healthy excretory channels, and must remain in the blood stream until subjected to proteolysis by the enzymes of the blood. In the course of ordinary life the quantities of such substances gaining entrance into the circulation are necessarily small, and would call forth but slight reactions. The sudden injection of large amounts of serum, not

⁹ Weichhardt, *Arch. f. Hyg.*, Vol. 74, 1911, has made similar studies and claims to have found toxic protein cleavage products similar to his kenotoxin in exposed air.

¹⁰ See Zinsser and Young, *Jour. Exp. Med.*, Vol. 17, 1913.

easily disposed of, would, on the basis of the preceding assumptions, result in a very gradual antigen destruction with consequent antibody formation, so that, at the end of eight to ten days, there would be present side by side remnants of unchanged antigen and newly formed specific antibody. The destroyed antigen fraction, in other words, gradually sensitizes the body to the fraction which persists and has not yet been assimilated or excreted at the end of this time. Such a point of view would explain, not only the reaction after a first injection, but would account for the incubation time in such cases, and for the differences between these reactions and both the "immediate" and the "accelerated" reactions of cases twice injected. The bearing which this point of view would have on the problems of incubation time in general is obvious.

The recognition of the anaphylactic nature of serum sickness has led to many attempts to develop methods of antitoxin administration by which these reactions could be avoided. Since it was determined that the degree of reaction was directly dependent upon the amount of the foreign serum injected, it was an obviously logical procedure to attempt in antitoxin production to concentrate as high a potency of antitoxin into as small as possible an amount of serum. Attempts have also been made to alter the serum itself in such a way that it would lose its properties of acting as an anaphylactic antigen without suffering materially in antitoxin value. Bujwid¹¹ found that serum sickness was less frequent after the use of sera which had been allowed to stand for prolonged periods, and we have seen that Besredka and others have claimed a reduction of toxic property in sera heated repeatedly to 60° C. It was hoped, moreover, that the so-called concentration methods—such as those of Gibson, Banzhaf, and others—would yield an antitoxin that would be devoid of anaphylactic properties. None of these methods of altering the serum can, however, be said to have been satisfactory in that the antitoxic property seems to be closely associated with the globulins,¹² which we have seen are at the same time closely associated with the production of anaphylaxis.

The conclusions of Rosenau and Anderson¹³ regarding this are based on direct experimentation with concentrated antitoxin made at the New York Department of Health by the Gibson method. They found the refined antitoxin, volume for volume, quite as toxic as the unrefined, but since the same amount of antitoxin is by this and other methods concentrated in a considerably smaller amount of

¹¹ Bujwid. Quoted from Friedberger and Mita, *Deutsche med. Woch.*, No. 5, 1912.

¹² Among others previously mentioned see also Turro and Gonzales, *C. R. de la Soc. de Biol.*, Vol. 69, 1910.

¹³ Rosenau and Anderson. *U. S. Pub. Health and M. H. S. Hyg. Lab. Bull.* 36, April, 1907.

protein solution there is a distinct gain for safety in the use of such preparations. Endeavors to produce potent antitoxic sera by chemical or physical methods without any sensitizing properties have thus been unsuccessful.

On the other hand, the knowledge gained by animal experimentation regarding the influence upon the anaphylactic manifestations exerted by various methods of administering the antigen has led to results which have proven of much value, both in the immunization of experimental animals and in human serum therapy. Probably the most carefully studied of these methods is the one which Besredka¹⁴ has recommended on the basis of his work on antianaphylaxis in animals. He found that sensitized guinea pigs could be injected with quantities of serum amounting to about one half or less of the fatal dose without showing symptoms, and subsequently, at intervals of 2 to 5 minutes, further injections of the serum could be given, the total amount five to twenty times exceeding the lethal dose without causing symptoms of any kind. From these experiments he has developed a method of serum injections the principle of which is very simply a division of dose. In lieu of injecting into an animal or man the entire quantity of serum at once, small, gradually increasing amounts are administered in two, three, or more doses, the intervals varying from five minutes to several hours, according to the necessities of speed indicated by clinical considerations. The process as applied to man consists, then, in preceding the injection of the larger quantity of the serum by one or two subcutaneous injections of smaller amounts. With this principle well defined it would be quite unwise to lay down definite rules of quantity or interval at present, since in no instance will it be possible to estimate the exact condition of susceptibility of the particular case. It goes without saying that the precautions should be particularly respected in children in whom the relation of 5 or 10 c. c. of serum volumes to the body weight approaches the dangerous proportions dealt with in animal experiments.

Besredka has also shown that if the rectum of a sensitive animal is cleaned out by enema, and a relatively large amount of the antigen then introduced, an injection may be given in within twelve to twenty-four hours later without danger, however delicate the hypersusceptibility of the animal has been. This method apparently must depend upon a slow, gradual absorption of antigen, and would seem to furnish a most convenient and advisable method to apply in man.

¹⁴ Besredka. *Ann. de l'Inst. Past.*, Vol. 24, 1910; *C. R. de la Soc. de Biol.*, 65, 1908, p. 478; *C. R. de la Soc. de Biol.*, Vol. 66, 1909, p. 125; *ibid.*, 67, 1909, p. 266; *C. R. de l'Acad. des Sc.*, Vol. 150, 1910, p. 1456; ref. *Bull. de l'Inst. Past.*, Vol. 8, 1910, p. 735.

Friedberger and Mita¹⁵ have suggested another method which also depends upon very slow administration rather than division of dose. In experiments upon guinea pigs they had found that sensitized animals which, as tested by controls, would succumb to intravenous injections of 0.01 c. c. of sheep serum per 100 grams weight when the entire quantity was injected within one minute, would survive a similar administration of as much as 0.1 c. c. if, by means of a specially constructed apparatus, the injection was made gradually, extending over a period of 100 minutes. While this method offers many practical difficulties to ordinary bedside application, it *does* show that the intervals of injections by the Besredka method do not need to exceed fractions of an hour—or, at most, a few hours—in order to add materially to the safety of injection.

There is another phase of specific therapy in which the question of possible anaphylaxis must be taken into consideration, and that is the treatment of patients with bacterial vaccines. As a matter of fact the danger of anaphylaxis in such cases is probably very remote—both because of the shortness of the intervals at which these injections are usually made and because of the extremely small amounts of protein represented by the usual dose of 100 or 200 millions of bacteria. However, the possibility cannot be disregarded, especially in children, and two cases were verbally described to the writer by Dr. Philip Van Ingen, in which gonococcus vaccines caused immediate symptoms of such a character that anaphylaxis could not be excluded.

Ohlmacher¹⁶ also has described localized reactions at the place of inoculation as well as swelling and tenderness at points of former inoculations following bacterial vaccine injections. He has occasionally seen slight systemic symptoms (dizziness, nausea, etc.) which he explains on the basis of anaphylaxis.

Moreover, it must be remembered that active sensitization with bacterial antigens has been most regularly successful in the hands of Kraus and Doerr,¹⁷ Holobut,¹⁸ and Kraus and Admiradzibi,¹⁹ as well as in confirmatory experiments carried out in the Stanford laboratory, when repeated injections at short intervals were made, rather than when, as in serum anaphylaxis, a single injection only was given. This would lend an even closer analogy to the procedures carried out during vaccine treatment. For instance, in the successful experiments of the last-named writers ten daily injections of 1/100 of a slant culture of dead colon bacilli were made for the purpose of

¹⁵ Friedberger and Mita. *Deutsche med. Woch.*, No. 5, 1912; figures taken from *Versuch.*, 3.

¹⁶ Ohlmacher. *Jour. Med. Res.*, Vol. 19, 1908, p. 113.

¹⁷ Kraus u. Doerr. *Wien. klin. Woch.*, No. 28, 1908.

¹⁸ Holobut. *Zeitschr. f. Immunitätsforsch.*, Vol. 3, 1909.

¹⁹ Kraus u. Admiradzibi. *Zeitschr. f. Immunitätsforsch.*, Vol. 4, 1910.

sensitization—the toxic dose of 1/2 slant being given fifteen days after the last of these.

In order to obtain some opinion regarding the possible dangers of vaccine therapy in this regard the writer a few years ago observed carefully a pair of young goats, animals extremely favorable for anaphylactic experiment (and of approximately the weight of a child of three) in the course of frequent and irregularly spaced intravenous injections of typhoid bacilli. In both cases marked anaphylactic symptoms were observed after the animals had attained a considerable agglutinative and bactericidal power (1 to 5,000 to 1 to 20,000), but in each case only after intravenous injections of large quantities of bacteria, 1/2 to 2 slant cultures. While, of course, such experiments are not conclusive in any way, from these, as well as from a number of laboratory accidents in the course of animal immunization, it is the writer's impression that the intravenous injection of bacteria or bacterial products in human beings would be a procedure involving some risk, unless more thorough experimental data than we at present possess were available to guide us as to dosage and intervals. The ordinary subcutaneous treatment of patients, however, with bacteria in the amounts customarily employed in vaccines would seem to be practically without risk as far as acute anaphylaxis is concerned.

In the treatment of animals with vaccines of various kinds Leclainche^{20 21} has repeatedly called attention to the fact that inoculation with a vaccine may lead to a condition of hypersusceptibility, serving to light up a latent lesion which might have been held in check if the normal resistance had not been interfered with. This objection, we have seen, has been made on numerous occasions against tuberculin therapy, and is one of the factors which have led to the great caution in dosage and control of all therapy based on active immunization. These considerations, even more than the rather remote dangers of serious active anaphylaxis, require that all forms of specific therapy should be carried out only under the safeguards of thorough familiarity with the experimental phases of such work.

Our own recent studies on anaphylatoxins, moreover, have inclined us to believe that hypersusceptibility to bacterial protein may well be a strong predisposing factor in infection.

Serum sickness, occurring as a direct consequence of the injection of a foreign protein into a human being, forces itself upon us as manifestly related to anaphylaxis. There are a number of other clinical conditions which are less obviously anaphylactic in nature, but in which we have many good reasons for attributing an important part of the etiology to a state of hypersusceptibility. Thus the pe-

²⁰ Leclainche and Vallée. *Ann. de l'Inst. Past.*, 1902.

²¹ Leclainche. *Revue Gén. Méd. Vét.*, Sept., 1911; *Bull. de l'Inst. Past.*, 9, 1911, p. 1089.

cular so-called "idiosyncrasies" observed in many people who suffer from urticarial skin rashes, gastro-intestinal difficulties, and even severe systemic illnesses after certain varieties of food seem to depend upon an acquired or possibly inherited hypersusceptibility to the particular proteins involved, which, at certain times of abnormal gastro-enteric conditions, can get into the circulation in small quantity. It is not impossible, furthermore, that such unfortunate cases as the severe forms of angioneurotic edema, which seem, at least in part, to be associated with gastro-intestinal disturbance, and which may be transmitted from mother to child, have their root in anaphylaxis. For this, however, we have only inference based on clinical observation.

ASTHMA AND HAY FEVER

Conditions in which there seems to be more definite ground for association with anaphylaxis are ASTHMA and HAY FEVER. In asthma the analogy has been clearly set forth by Meltzer.²² He points out that in both asthma and anaphylaxis the symptoms consist in a tonic stenosis of the small bronchioli of peripheral origin, and that both conditions are favorably affected by the administration of atropin. It is, of course, not certain, but it seems extremely likely that so-called "nervous asthma" is nothing else than an anaphylactic attack in a hypersusceptible individual when the particular protein to which he is sensitive gains access either by the alimentary or respiratory path.

Very closely related to asthma is the condition known as "hay fever." This disease has been of recent years most thoroughly studied by Dunbar.²³ Dunbar has ascertained that the hay fever prevalent in Europe is dependent chiefly upon a protein substance found in the pollen of most grasses, while that of America, which occurs chiefly in the autumn, is caused by the proteins of the pollen cells of the ambrosiaceæ and solidagineæ—plants which are generally distributed on the North American continent and bloom in August and September. The disease occurring in China is caused by another plant, the *Ligustrum vulgare*. The suggestion that the disease was due to anaphylactic action of these pollen proteins upon hypersusceptible individuals was first made by Wolff-Eisner.²⁴ Dunbar has gone into the question with great thoroughness, and has come to the conclusion that the disease has much in common with anaphylaxis—though he believes that, in addition to a hypersusceptibility to the pollen "toxin," there must be present in the patients, at the same

²² Meltzer. *Jour. of the A. M. A.*, Vol. 55, 1910, p. 1021.

²³ Dunbar. *Berl. klin. Woch.*, Nos. 26, 28, 30, 1905; *Zeitschr. f. Immunitätsforsch.*, Vol. 7, 1907; *Deutsche med. Woch.*, Vol. 37, 1911, p. 578.

²⁴ Wolff-Eisner. "Das Heufieber sein wesen u. seine Behandlung," 1906.

time, an abnormal "Durchlässigkeit" or penetrability of the cutis and mucosa for the pollen substances. He claims to have shown that a solution of pollen protein instilled into the eye—or even dropped upon the skin of a hay-fever patient—gives rise to a prompt and severe reaction, while it produces no effect upon normal persons. Unlike experimental serum anaphylaxis, the repeated instillation of the pollen substances rather increases than diminishes the susceptibility even when these are carried out daily. Furthermore, unlike serum anaphylaxis, against the manifestations of which no direct passive immunization has so far been possible, Dunbar claims to have produced a curative immune serum by the treatment of horses with the pollen extracts ("Pollantin"). Dunbar, therefore, while admitting an anaphylaxis-like hypersusceptibility of the patients, still believes that the antigen in this case is a true "toxin" against which an antitoxin can be produced—the condition being more directly comparable to the sensitization against diphtheria and tetanus toxins observed during the earlier phases of these investigations by v. Behring and his associates rather than to the phenomena of serum anaphylaxis themselves.

Schittenhelm and Weichhardt,²⁵ on the other hand, regard hay fever as truly anaphylactic in every sense. They speak of it as epithelial anaphylaxis (hay fever being specifically designated as "conjunctivitis and rhinitis anaphylactica," in distinction from other forms of cellular anaphylaxis, i. e., enteritis anaphylactica). They believe that the manifestations of the disease result from a local hypersusceptibility in which a toxic substance (Abbau Produkt—similar to anaphylatoxin) is produced. The so-called "antitoxin" of Dunbar acts favorably only when locally applied, and not on subcutaneous administration. For this reason they do not regard it as a true antitoxin, but think it acts as a local antiferment which prevents or delays the cleavage of the pollen-substance into its toxic split-product—thereby preventing or ameliorating the attacks.

Similar to hay fever are the sudden attacks of catarrhal nasopharyngitis and conjunctivitis—often of asthma-like respiratory difficulty, with itching of the nose and eyes and sneezing which many individuals experience when coming close to horses, cats, or other animals. In the Stanford University laboratory the writer had an assistant who invariably had such attacks, sudden, violent, and of several hours' duration, when handling guinea pigs for experiment. The character of such attacks has long aroused the suspicion that the reaction was anaphylactic in nature, especially since it was known that extremely slight amounts of antigen could give rise to symptoms in susceptible subjects. The difficulty in these cases was the question of the nature of the antigen which emanated from the animal to excite an attack. Recently, however, observations having impor-

²⁵ Schittenhelm and Weichhardt. *Deutsche med. Woch.*, 37, No. 19, 1911.

tant bearing upon this problem have been made by Weichhardt²⁶ and by Rosenau,²⁷ who have demonstrated the presence of organic matter in expired breath. Rosenau condensed the moisture of the expired breath of man and injected the liquid so obtained into guinea pigs. After two weeks these animals were injected with normal human serum, and out of 99 test animals 26 responded with symptoms of anaphylaxis. This demonstrated not only the presence of organic matter in the breath, but showed at the same time that such organic matter was probably protein in nature or at least surely capable of acting as anaphylactic antigen. Rosenau surmises, therefore, that such protein may be slightly volatile under the given conditions. He suggests that sensitization in this manner may explain the harmful effects resulting from a first injection of horse serum into patients, previous sensitization having occurred by close association with horses. Surely it would explain logically the "cellular" or epithelial anaphylaxis experienced by certain people in the presence of animals. In our opinion this is rendered more likely, since, in the case mentioned as occurring at Stanford University, the inhalation of washings (both aqueous and alcohol soluble) obtained from the hair and skin of guinea pigs, and dried in Petri dishes, produced absolutely no effects in the susceptible individual, whereas continued handling of a living pig almost invariably caused such marked effects that the person in question often became useless as an assistant because of violent attacks of sneezing. It must not be omitted, however, that not all observers have confirmed Rosenau's work, and his explanation must therefore be regarded as merely tentative.

An interesting train of suggestions connecting human pathology with anaphylaxis has followed the discovery of "organ-specificity" in the case of hypersusceptibility similar to that noted by Uhlenhuth in connection with the precipitin formation and described in another chapter.

It was shown by Kraus, Doerr, and Sohma,²⁸ we have seen, that animals sensitized with the crystalline lens protein of one animal species would react to lens protein in general, and not necessarily to the tissue protein of the animal species from which it was taken. In other words, the ordinary "species" specificity did not hold good. Specificity was determined in this case by the character of the organ rather than by that of the species. The same thing was shown for testicular protein by v. Dungern and Hirschfeld.²⁹ The proteins of these organs from various animals have therefore a certain common antigenic property which is independent of the antigenic element

²⁶ Weichhardt. *Arch. f. Hyg.*, Vol. 74, 1911.

²⁷ Rosenau and Amoss. *Jour. Med. Res.*, Vol. 25, Sept., 1911.

²⁸ Kraus, Doerr, and Sohma. *Wien. klin. Woch.*, Vol. 21, 1908, p. 1084.

²⁹ Von Dungern u. Hirschfeld. *Zeitschr. f. Immunitätsforsch.*, 4, 1910.

common to the particular species. Further than this, Andrejew³⁰ claims to have shown that it is possible to sensitize an animal with its own lens protein. A few guinea pigs injected by him with their own lens proteins, and reinjected with the same substances after a suitable interval, reacted with definite anaphylactic symptoms. The possibility is thus given that an animal or human being could become sensitized by its own organ proteins if these were traumatically or otherwise destroyed and absorbed. The train of reasoning is similar to that which has given much hope of enlightenment to pathologists when the earlier work upon the cytotoxins was done. Rosenau and Anderson,³¹ for instance, injected guinea pigs with guinea pig placenta, and found that, after the usual period of incubation, the animals reacted to a second injection with marked symptoms of anaphylaxis. On the basis of these experiments Rosenau and Anderson suggest that certain of the toxemias of pregnancy are of anaphylactic origin. They believe that it is possible that a mother may become sensitized by the "autolytic products of her own placenta," the result being eclampsia.

By a similar process of reasoning Elschnig³² has attempted to explain sympathetic ophthalmia. He claims to have shown that the laws of organ specificity apply to the proteins (especially the pigment) of the uveal tract. The destruction and absorption of injured uveal tissue, according to him, induce the formation of organ-specific antibodies by which the remaining uveal structures of the same, as well as of the opposite, eye are sensitized. The consequence is a "sympathetic" inflammation which "is to be regarded purely as an anaphylactic reaction."

These and other similar suggestions less well founded experimentally illustrate the possibilities for clinical reasoning furnished by a knowledge of the anaphylactic phenomena. In no cases of this sort, however, can the association with anaphylaxis be as yet regarded as more than an extremely interesting suggestion.

From all that has gone before it is quite evident that most of the positive facts which may be regarded as determined concerning the phenomena of anaphylaxis have been obtained in experiments with small and very sensitive animals, comparatively large and measured quantities of antigen, and often by the violent method of intravenous injection in which the entire mass of antigen comes rapidly into contact with the available antibodies and the vulnerable tissues. We cannot, therefore, draw rigid parallels between these experiments and clinical manifestations in human beings in whom

³⁰ Andrejew. *Arb. a. d. kais. Gesundh.*, Vol. 30, 1909.

³¹ Rosenau and Anderson. *U. S. Pub. Health and M. H. S. Hyg. Lab. Bull.* 45, 1908.

³² Elschnig. *Von Graefe's Archiv f. Ophthal.*, Vol. 75, p. 459; Vol. 76, p. 509; Vol. 78, p. 549.

the localization, quantitative discharge of antigen, and consequent production of antibodies is of necessity irregular and different in each individual case. We have learned, as a general conception, however, that the introduction into the animal body of antigenic substances of all varieties leads, under certain conditions, to increased tolerance or resistance—under other circumstances to a state of greater susceptibility—these diametrically opposed physiological consequences being, in all probability, determined by relative concentrations of antigen and antibody, their speed of contact, and their quantitative relationship to available alexin. The problems of clinical medicine in which the possibility of anaphylaxis can be at all considered, therefore, are extremely complicated, and few of them can be approached by direct experiment.

In the case of serum sickness the analogy has been so clear and the experience with human beings so extensive that practically no doubt can exist as to the common mechanism of this condition with that of experimental anaphylaxis. In the other conditions mentioned the connection is one of great likelihood, but after all is inferential, and calls for much further investigation. For this reason it is best to abstain from a further enumeration of many other maladies in which the condition of hypersusceptibility has been suggested as a vaguely possible etiological factor.

ANAPHYLAXIS AND THE TUBERCULIN REACTION

There is one class of phenomena, however, which calls for further discussion in this connection, since its dependence upon anaphylaxis, while generally assumed, is still opposed by many authorities. This consists of the various DIAGNOSTIC REACTIONS in which extracts of micro-organisms are injected, or brought into contact with the skin or conjunctiva of infected subjects. Such are the various forms of the *tuberculin reaction*, the *typhoid reaction* of Chantemesse, the one of Gay, and the *luetin reaction* of Noguchi. In the tuberculin reaction the conditions have been thoroughly studied, and we may make a detailed consideration of this example serve to bring out the general principles involved.

In all forms of the TUBERCULIN REACTION there is a very evident hypersusceptibility to various forms of antigen derived from the bacillus. When the tuberculin is injected subcutaneously the reaction is systemic and also localized to a certain extent in any tuberculous foci which may be present. When the v. Pirquet or Moro skin reactions are carried out, or the Calmette ophthalmic test is made, the reactions are almost purely local. In all cases reactions are induced by quantities of antigen which cause no effect whatever in normal individuals.

The basic observation leading to the diagnostic use of tuberculin was made by Koch³³ upon guinea pigs. He describes his observation as follows:

"Tuberculin may be injected into normal guinea pigs in considerable quantities without causing noticeable symptoms. Tuberculous guinea pigs, on the other hand, react to comparatively small doses in a very characteristic manner."

Since, in Koch's experiments upon tuberculin, it was desirable for his particular purposes at the time, to obtain very sharp reactions, he did not content himself with the production of moderate symptoms by the injection of slight amounts of tuberculin into infected animals, but increased his dosage until the guinea pigs were killed. He showed that guinea pigs having a moderately advanced infection—4 to 5 weeks after inoculation—could be killed by doses of 0.2 to 0.3 gram, while animals in very advanced stages would succumb within 6 to 30 hours to quantities as small as 0.1 gram subcutaneously. In the animals so studied he determined not only a systemic effect, but a very marked local reaction as well in the skin, areolar tissues, and adjacent lymph nodes.

Koch's observations upon guinea pigs were applied by him, Guttstadt,³⁴ Beck,³⁵ and others to man, and the result was the development of the present important diagnostic test. The fundamental fact in this as well as in other tests of this kind, then, is the appearance of local and systemic reactions in infected subjects to contact with specific antigenic material which, at least in the same quantities, produces no effects in normal individuals. The analogy with the phenomena of anaphylaxis is thus indicated.

Koch's original interpretation of the phenomenon was of course unaided by any of the later observations on anaphylaxis. According to him the tuberculin contained substances which caused tissue necrosis. The necrotizing action was particularly powerful upon tissues which were tuberculous, and therefore already saturated with the toxic material. The destruction of such tissues resulted in systemic symptoms.

Very similar to this view is the one later expressed by Babes and Broca,³⁶ who attribute the systemic symptoms to a sudden lighting up of the existing lesions by the small amount of extra tuberculin added to that already present in these foci.

The first suggestion of the possible association of the tuberculin reaction with the union of an antigen and its antibody was made by Wassermann and Bruck.³⁷ They accepted Ehrlich's assumption

³³ Koch. *Deutsche med. Woch.*, No. 43, 1891.

³⁴ Guttstadt. "Klin. Jahrbuch" *Ergänzungsband*, 1891.

³⁵ Beck. *Deutsche med. Woch.*, No. 9, 1899.

³⁶ Babes u. Broca. *Zeitschr. f. Hyg.*, Vol. 23, 1896.

³⁷ Wassermann and Bruck. *Deutsche med. Woch.*, No. 12, 1906.

that certain cells of the tuberculous foci (those situated just below the periphery and already affected by the tubercle toxin, though still resistant) were possessed of an increased receptor apparatus for the tubercle antigen. For this reason the injected tuberculin was concentrated in these foci, attracted out of the circulation by the increased avidity of these cells, the consequence being increased activity of the lesions and systemic symptoms. The tuberculin reaction, according to these writers, therefore, would be caused by the union of the tuberculin with the "sessile receptors" upon the diseased tissues—a point of view which would specify the diseased tissues and their products as the sources from which emanated the toxic factors inciting the systemic symptoms.

The theories of Koch and of Babes do not, as Meyer points out, explain the frequent absence of the tuberculin reaction in very advanced cases of human tuberculosis, as contrasted with its frequency and regularity in the earlier cases. For, according to both of these views, the more severe the existing lesions the more actively would the injected tuberculin initiate tissue necrosis and consequent symptoms. The theory of Wassermann and Bruck avoids this objection since it presupposes the acceptance of Ehrlich's view that the increased receptor apparatus is present and free only in those cells in which necrotic destruction has not yet set in. In the necrotic areas the receptor apparatus is already saturated or satisfied as to its affinities, and extensive areas of necrosis, therefore, are unaffected by contact with further quantities of tuberculin.

All of these theories, however, inasmuch as they refer the tuberculin reaction to alterations taking place in more or less active lesions, are unable to account for the occurrence of the reaction in persons in whom healed foci only are present, and are entirely inconsistent with the facts we now possess regarding the cutaneous and ophthalmic tests in which the reactions occur in previously healthy tissues.

These facts practically exclude the acceptance of any theories which regard the tuberculous focus as the sole source of the reaction. We may still accept the Koch or Wassermann views to explain local swellings and other changes in infected lymph-nodes or other lesions, but we must assume in addition to this a generalized hypersusceptibility at least analogous to the phenomena of anaphylaxis.

This ability of previously healthy tissues, remote from any center of tuberculous infection, to react to the application of tuberculin was discovered by von Pirquet³⁸ in the development of his skin reaction, and by Calmette³⁹ and Wolff-Eisner⁴⁰ in their work upon

³⁸ v. Pirquet. *Berl. klin. Woch.*, No. 20, p. 644, and No. 22, p. 699, 1907; also "Klinische Studien über Vaccination," Deuticke, Wien, 1907.

³⁹ Calmette. *C. R. de l'Acad. des Sciences*, June, 1907.

⁴⁰ Wolff-Eisner. *Berl. klin. Woch.*, 1907, p. 1052. Discussion of paper by Citron.

the ophthalmoreaction. The principle involved in these reactions was then further emphasized by the introduction of the Moro tuberculin-ointment method and the intracutaneous test of Römec. The mere observation that the infection with tuberculosis results in a general tissue hypersusceptibility immediately suggests the interpretation of the tuberculin reaction as a manifestation of anaphylaxis. Von Pirquet, accordingly, on the basis of his previous studies upon serum sickness includes the reaction in the category of what he calls "allergie."

He assumes that the reaction depends upon the presence in the system of antibodies, which form a union with the applied tuberculin, the result being the formation of poisons and a reaction. This assumption, according to the anaphylatoxin theory of Friedberger, would imply the participation of alexin in the reaction—acting upon the united tuberculin-antituberculin complex, though v. Pirquet does not express himself positive as to this.

This, moreover, is the clearly expressed opinion of Friedberger⁴¹ himself. Consistently with his general theory of anaphylaxis he assumes that the injected tuberculin comes into relation with specific antibodies with which it unites, the alexin then splitting off anaphylatoxin from the complex. He bases this view upon his experimental demonstration, mentioned above, of the production of anaphylatoxin from tubercle bacilli by *in vitro* digestion with guinea pig complement.

In principle the view of v. Pirquet is similar to that previously expressed by Wolff-Eisner⁴² that the union of tuberculin with its lytic antibody, present in the tuberculous animal, gave rise to poisons as the result of lysis. Both of these theories simply apply to the special case of the tuberculin reaction theories of mechanism applied to anaphylactic reactions in general.

We must admit that the facts of the "allergie" reactions as a class seem to force upon us the acceptance of von Pirquet's views. Apart from the purely clinical observations made in carrying out the routine tests we have the additional evidence that the instillation of tuberculin into the eye of normal individuals gives rise to no reaction, but a repetition of the instillation into the same eye after ten or more days results in a marked and typically positive test. Furthermore, von Pirquet⁴³ states that individuals showing no clinical tuberculosis and negative to a first test will often react ("sekundäre Reaktion") to a second test carried out a few days after the first.

These facts all seem to indicate acquired hypersusceptibility more analogous to true serum-anaphylaxis than to the toxin hypersuscepti-

⁴¹ Friedberger. *Münch. med. Woch.*, Nos. 50 and 51, 1910.

⁴² Wolff-Eisner. *Berl. klin. Woch.*, Nos. 42 and 44, 1904.

⁴³ Cited from Löwenstein in "Kraus u. Levaditi Handbuch," Vol. 1, p. 1039.

bility of von Behring, in that the tuberculin is but slightly toxic in itself.

If the analogy is such a close one, therefore, it should be easy to formulate experiments by which the phenomena now ascertained regarding serum-anaphylaxis could be demonstrated for tuberculin hypersusceptibility. The obvious procedure, therefore, would be to attempt to passively transfer tuberculin sensitiveness to a normal animal with the serum of a tuberculous one. This has indeed been attempted by Friedemann,⁴⁴ later by Bauer⁴⁵ and a number of others—usually with negative result. The writer, hoping to develop a diagnostic method for tuberculosis, has also attempted this by the transference of human tuberculous blood to guinea pigs, but invariably obtained negative results. Yamanouchi⁴⁶ alone has reported positive experiments by a similar procedure with rabbits, but so far his results, according to Friedemann, have completely failed of confirmation. Austrian succeeded by sensitizing guinea pigs with 5 c. c. of titrated whole blood, using for the second injection a tuberculo-protein prepared by the method of Baldwin.⁴⁷ In this particular, therefore, the analogy between anaphylaxis and the tuberculin reaction, though not easily worked out, has nevertheless been established. Another objection which has been made by a number of observers is the fact that anaphylaxis is accompanied by temperature depression while tuberculin reactions are followed by a rise. This objection may be regarded as invalid, however, in the light of Friedberger's⁴⁸ experiments which showed that temperature depression follows only when large doses of the antigen are injected into the sensitized animals, smaller doses often giving rise to increased temperature.

We gain a certain amount of insight into the conditions here prevailing by considering the information which has been obtained from the study of the antibodies formed in animals in tuberculosis. It appears from the work of Christian and Rosenblatt⁴⁹ that antibodies to the tubercle bacillus are formed by tuberculous animals only. Normal animals form these to a very slight degree only, if at all, when immunization with tuberculin is attempted. In other words, as Friedemann ("Weichhardt's Jahresbericht," 6, 1910) points out, the specific reaction of antibody formation in tuberculosis seems to be closely associated with the tuberculous tissues themselves.

⁴⁴ Friedemann. "Über anaphylaxie," "Weichhardt's Jahresbericht," Vol. 6, 1910.

⁴⁵ Bauer. Cited *ibid.*; also *Münch. med. Woch.*, 1909, p. 1218.

⁴⁶ Yamanouchi. *Wien. klin. Woch.*, 1908, p. 1263.

⁴⁷ Austrian *Bull. of the Johns Hop. Hosp.*, Vol. 24, 1913; Baldwin, *Journ. Med. Res.*, Vol. 17, 1910.

⁴⁸ Friedberger. *Deutsche med. Woch.*, No. 11, 1911.

⁴⁹ Christian and Rosenblatt. *Münch. med. Woch.*, 1908.

The same inference can be made from Bail's⁵⁰ experiments on passive sensitization. For, although passive sensitization of guinea pigs with the serum of tuberculous animals has been unsuccessful, Bail succeeded in obtaining lethal anaphylactic reactions by injected macerated tuberculous tissues, following these on the next day by injections of tuberculin. It is plain from this, as Friedemann correctly argues, that we must assume that the antibodies (receptors) formed against tubercle bacilli are closely bound up with the tissue cells, the reaction of tuberculin being largely with "sessile receptors." Indeed, it seems as though the antibodies formed against tubercle bacilli undoubtedly remain in close relation to the cells of the tissues except in cases of active tuberculosis in which localized areas of cells are under the influence of a very intense action of the poisons, and a consequent overproduction and discharge of receptors (using the Ehrlich nomenclature) may occur. This would correspond with considerable accuracy, moreover, to the histological facts, for in this infection, similar to leprosy and a number of other conditions, but unlike most acute infections, the battle against the micro-organisms is carried out chiefly by the adjacent tissue cells.

We might assume, therefore, that, in tuberculous individuals, there is indeed a reaction, at first local, then generalized to a slight degree, in which antibodies are actually formed. These antibodies, however, remain to a preponderant extent sessile, or incorporated in the reacting cells. Upon the injection or application of tuberculin the reaction takes place in or upon the cells. Whether or not the further coöperation of complement or alexin is then necessary for the lysis and poison production from the antigen, as in the similar reactions taking place in the circulation, or whether the intracellular ferments themselves suffice for this, cannot be decided at present. It is certainly not unlikely that the circulation of tubercle-antigen—even in small quantities—throughout the body may produce such hypersusceptibility of cells (represented graphically by the conception of sessile receptors in many parts of the body remote from the lesion—a quality remaining constant for prolonged periods and explaining the subsequent skin and ophthalmic reactions obtained upon test). Certain clinical observations cited by v. Pirquet⁵¹ would seem to support such a view. For instance, he states that, having employed his left forearm repeatedly for tests, he was able to obtain positive reactions in this area with tuberculin diluted 1 to 1,000, whereas his right forearm was negative to tuberculin ten times as concentrated. Furthermore, as Kohn⁵² has shown that, while the first injection of tuberculin into the eye of a normal person produces no reaction, this eye will not only react to a second instillation,

⁵⁰ Bail. *Zeitschr. f. Immunitätsforsch.*, Vol. 4.

⁵¹ V. Pirquet, "Kraus u. Levaditi Handbuch," Vol. 1, p. 1050.

⁵² Kohn. Quoted from Löwenstein, Kraus and Levaditi, Vol. 1, p. 1033.

but will show a reaction when the second application is made subcutaneously. Negative evidence pointing in the same direction is the observation that absolutely no influence is exerted upon the outcome of tuberculin reaction if the tuberculin is previously mixed with blood serum from either positively or negatively reacting cases.⁵³ This would tend to show that, whether reacting or not, the factors which determined this are certainly not present in the circulating plasma.

That the circulation of tubercle antibodies in the blood may even interfere with the localized tuberculin reaction is rendered likely by the fact that skin reactions are often negative in cases of advanced tuberculosis, and that, as we are told by Dr. Blair, of the N. Y. Zoölogical Park, such reactions are usually negative in tuberculous monkeys in which the disease is invariably very rapid and acute.

PRACTICAL DIAGNOSTIC USES OF ANAPHYLAXIS

The specificity of the anaphylactic reaction has led to extensive attempt to utilize it for forensic protein determinations in the same way in which the precipitin test is used. Uhlenhuth,⁵⁴ Thomsen,⁵⁵ Pfeiffer, and others have carried on extensive experimentation in this problem, the technique, in general, consisting in sensitizing guinea pigs with solutions of the unknown protein (dissolved blood spots, etc.) and testing them by a second injection of the suspected protein after the usual anaphylactic incubation time. The results of such work have shown that indeed positive reaction may be obtained and diagnosis made in this way. However, the reactions are not ordinarily very striking, and this method is not as reliable as the precipitin method. Uhlenhuth⁵⁶ believes that the anaphylactic reaction has value only in cases in which the amount of unknown protein is so small or so changed by preservation or decomposition that its precipitable qualities have been lost.

Yamanouchi's⁵⁷ attempt to utilize anaphylaxis for the diagnosis of tuberculosis, by passively sensitizing guinea pigs with the serum of tuberculous patients and testing subsequently with tuberculin, has been mentioned before. Although he claims positive experiments, our own experience with a similar technique has given us results which were so irregular that we feel that this technique has very slight practical value, if any.

Pfeiffer⁵⁸ has attempted to apply anaphylaxis to the diagnosis

⁵³ V. Pirquet. *Loc. cit.*

⁵⁴ Uhlenhuth. *Deutsche milit. Zeitschr.* No. 2, 1909. Cited from same author, *Zeitschr. f. Immunitätsforsch.*, Vol. 1, 1909.

⁵⁵ Thomsen. *Zeitschr. f. Immunitätsforsch.*, Vol. 1, 1909.

⁵⁶ Uhlenhuth. *Zeitschr. f. Immunitätsforsch.*, Ref. Vol. 1, 1909, p. 525.

⁵⁷ Yamanouchi. *Wien. klin. Woch.*, No. 47, 1908.

⁵⁸ Pfeiffer. *Zeitschr. f. Imm.*, Vol. 4, 1910.

of malignant tumors. Together with Finsterer⁵⁹ he sensitized guinea pigs with the serum of carcinomatous patients following the injection 48 hours later with press-juices of tumors. His conclusions were drawn from the anaphylactic temperature reaction, and he claims that animals so sensitized are hypersusceptible to the juices obtained from carcinomata, whereas animals sensitized with normal serum or the serum of sarcoma patients show no hypersusceptibility. Ranzi⁶⁰ has not been able to confirm this. The significance of such experiments, if correct, apart from their practical value for the diagnosis of carcinoma, would be considerable in that they tend to show that cancer tissues contain a specific protein which is antigenically distinct from the other tissue-proteins of the afflicted individual. However, we cannot yet accept these facts as absolutely established.

⁵⁹ Pfeiffer and Finsterer. *Wien. klin. Woch.*, No. 28, 1909.

⁶⁰ Ranzi. *Zeitschr. f. Imm.*, Vol. 2, 1909.

CHAPTER XIX

THERAPEUTIC IMMUNIZATION IN MAN

FACTS CONCERNING ANTITOXIN TREATMENT IN MAN

THERAPEUTIC USE OF DIPHTHERIA ANTITOXIN

IT is not consistent with the purpose of this brief treatise to discuss extensively the therapeutic benefits obtained by serum therapy in diphtheria. We can convey briefly an adequate idea of this by citing some of the tables given by Northrup in Nothnagel's "Encyclopedia of Practical Medicine," American Edition, Volume on Diphtheria, etc., p. 143. These figures are taken from the statistics of the New York Board of Health, which began treatment of diphtheria with antitoxin in January, 1895. Dr. Northrup states, however, that serum treatment cannot be considered to have been in general use until some time later.

Without Antitoxin

| Year | Cases reported | Deaths | Mortality, per cent. |
|------------|----------------|--------|----------------------|
| 1891..... | 5,364 | 1,970 | 36.7 |
| 1892..... | 5,184 | 2,106 | 40.0 |
| 1893..... | 7,021 | 2,558 | 36.4 |
| 1894..... | 9,641 | 2,870 | 29.7 |
| Total..... | 27,210 | 9,504 | Avg. 34.9 |

With Antitoxin

| | | | |
|------------|--------|-------|-----------|
| 1895..... | 10,353 | 1,976 | 19.0 |
| 1896..... | 11,399 | 1,763 | 15.5 |
| 1897..... | 10,896 | 1,590 | 14.5 |
| 1898..... | 7,173 | 919 | 12.8 |
| 1899..... | 8,240 | 1,085 | 13.1 |
| 1900..... | 8,364 | 1,176 | 14.0 |
| Total..... | 56,425 | 8,509 | Avg. 15.0 |

Table taken directly from Northrup, *loc. cit.*

From this table there appears a reduction of 58 per cent. in mortality and a similar drop is evident from the German statistics of Dieudonné,¹ from those of Welch, and many others.

It should be considered, moreover, in reading such statistics that they are made on gross mortality reports without elimination of the many cases that have not come under observation until too severely diseased to react to any form of treatment. The reason for the failure to obtain results with antitoxin when the cases have proceeded beyond a certain stage of intoxication will become evident when we consider the manner of absorption of the poison in a succeeding paragraph. The mortality sinks to between 8 and 9 per cent., when such cases are omitted, as is shown by the collective investigations of the American Pediatric Society in 1896—figures which we take also from Northrup's comprehensive study. This purely statistical evidence, however good, is further reënforced by the unquestionable and considerable diminution of emergency operations,² such as intubation and tracheotomy, since introduction of the antitoxin. Moreover, there is the manifold clinical evidence of benefit, after the serum treatment, familiar to every practicing physician.

Although the injection of antitoxin is of benefit by whatever route and in whatever quantity it may be given, nevertheless recent experimental investigations have taught us much regarding the proper use of this therapeutic agent. Especially interesting are the investigations of Meyer,³ who showed the extreme importance of an early use of the antitoxin. Apparently, as we have mentioned in another place, like tetanus antitoxin, the diphtheria poison may be in part absorbed directly by the nerves.⁴

There is apparently a great difference in therapeutic efficiency, according to the method in which the serum is administered, a difference probably depending upon speed of absorption. Berghaus⁵ showed that intravenous injection is 500 times more potent therapeutically than the subcutaneous, and 80 to 90 times more so than the intraperitoneal injection. Schick, in discussing this problem from the clinical point of view, for this reason lays special stress upon the speed of administration. He says: "Not only days but hours are of great importance." He bases this opinion largely upon the fact that the toxin which has already united with the nerve substance can probably no longer be influenced by the injection of the serum.

According to the experiments of Meyer and Ranson diphtheritic

¹ Dieudonné. *Arb. aus dem kais. Gesund.*, XIII, 1897.

² Siegert. "Jahrbuch f. Kinderheilkunde," Vol. 52, cited after Wernicke.

³ Meyer. *Berl. kl. Woch.*, 25, 26, 1909; *Arch. f. exp. Path. u. Ther.*, Vol. 60, 1909, and *Berl. kl. Woch.*, No. 45, 1911.

⁴ For a thorough discussion of these conditions see Schick, *Centralbl. f. Bakt.*, Rev. Vol. 57, 1913, "Report of 7th Meeting of the Mikrobiol. Gesell.," Berlin, 1913.

⁵ Berghaus. Cited from Schick, *loc. cit.*

paralysis may follow even when vigorous serum treatment has been employed. For, according to them, only the toxin which has reached the central nervous system through the circulation can be influenced by the serum, but no effect is possible upon the fraction which has been absorbed from the nerve endings directly.

Schick,⁶ on the basis of extensive experiments, comes to the conclusion that the subcutaneous injection of 1,000 to 2,000 units in diphtheritic cases has an immunizing value only, which protects the tissues from further injury and leads to cure if, at the time of injection, the lethal dose has not yet united with the sensitive cells. "If," he states, "we wish to obtain antitoxic action upon toxin which has already gone into action before the injection of the serum, then results can be obtained both in man and in animals only if a great deal of antitoxin is injected intramuscularly or intravenously."⁷

Interesting also from a clinical point of view are the studies of Schick,⁸ Hahn,⁹ and others¹⁰ upon the presence of antitoxin in the blood of normal, untreated individuals at different ages. These investigations were carried out by the intracutaneous method of toxin and antitoxin determination described in greater detail in a later section. The following table, taken from the article of Hahn, illustrates the experience, in such investigations, both of Schick and of Hahn himself. The determinations were carried out upon individuals who had never had diphtheria, as far as could be learned.

| Age | | Cases with antitoxin serum | Cases without antitoxin serum | Highest antitoxin value in 1 c. c. ¹¹ |
|------------|-----------------|-------------------------------|----------------------------------|---|
| Schick.... | { Newborn.... | 11 | 0 | under 1.5 units |
| | { 0-1 year.... | 1 | 3 | 0.11 unit |
| | { 2-10 years.. | 7 | 5 | 1.0 unit |
| | { 11-20 years.. | 8 | 9 | 0.75 unit |
| Hahn..... | { 21-30 years.. | 9 | 5 | 2.5 units |
| | { 31-40 years.. | 5 | 1 | 0.25 unit |
| | { 41-65 years.. | 2 | 8 | 2.5 units |

The table shows that in newborn children there is almost regularly a definite and sufficient protective value in the serum which diminishes up to the first year, so that at the end of the first year three out of four individuals had no antitoxin in their serum. In subsequent years up to the age of 40 an increasing percentage of

⁶ Schick. *Loc. cit.*

⁷ Schick. *Loc. cit.*, p. 32.

⁸ Schick. "Über Diphtherimmunität," Wiesbaden, 1910.

⁹ Hahn. *Deutsche med. Woch.*, Vol. 38, No. 29, p. 1366, 1912.

¹⁰ Karasawa and Schick. *Zeitschr. f. Kinderkrankheiten*, 1910, and "Jahrbuch f. Kinderheilkunde," 1910.

¹¹ Table taken directly from Hahn, *loc. cit.*

people have sufficient amounts of diphtheria antitoxin in their blood. After the age of 40 an increasing percentage is without such protection. The first observation, that newborn children usually possess considerable amounts of antitoxin, is very probably due to passive immunization by the blood of the mother, a fact which we have mentioned in another place. The exact method by which such measurements are made is described in a subsequent section on the intracutaneous method of determining toxin and antitoxin.

The work of Schick, that of J. Henderson Smith, and recent studies by Park and Biggs promise to alter considerably the methods of antitoxin therapy as at present in use in diphtheria. Smith measured the speed of absorption of antitoxin injected subcutaneously into the abdominal wall of a healthy man. His results are shown in the following table, which we take from his communication (page 213):

TABLE V

One c. c. of the patient's serum contained:

| Before injection | No demonstrable antitoxin |
|-------------------------------|---|
| 5 hours after injection..... | 0.1 unit antitoxin |
| 14 hours after injection..... | 0.225 unit antitoxin |
| 32 hours after injection..... | 0.68 unit antitoxin |
| 44 hours after injection..... | 1.0 unit antitoxin |
| 3 days after injection..... | 1.3 units antitoxin |
| 4 days after injection..... | 1.3 units antitoxin |
| 6 days after injection..... | 0.68 unit antitoxin |
| 13 days after injection..... | 0.17 unit antitoxin |
| 15 days after injection..... | 0.14 unit antitoxin |
| 20 days after injection..... | 0.08 unit antitoxin |
| 27 days after injection..... | No demonstrable antitoxin ¹² |

Park and Biggs¹³ have made similar studies and have contrasted the speed of absorption after subcutaneous administration with that after intravenous injection, basing their curves upon careful measurements of the sera of the treated patients. We reproduce their charts as given in their recent publication.

It is apparent from these charts, as well as from the work of Henderson Smith, that antitoxin, subcutaneously given, is slowly absorbed, and does not reach its maximum concentration in the blood stream until forty-eight hours or more after the injection. It follows that, as Park and Biggs point out, it is more rational to inject a single adequate dose than to divide the dosage and inject at intervals. They have obtained results in animal experiment which graphically illustrate this principle. A rabbit which had received ten fatal doses of toxin intravenously was given a total of 500 antitoxin units in divided doses as follows: 100 units after twenty min-

¹² J. Henderson Smith. *Journal of Hygiene*, Vol. 7, 1907, p. 205.

¹³ Park and Biggs. *Collected Studies from the N. Y. Department of Health, Bureau of Laboratories*, Vol. 7, 1912-1913, p. 27.

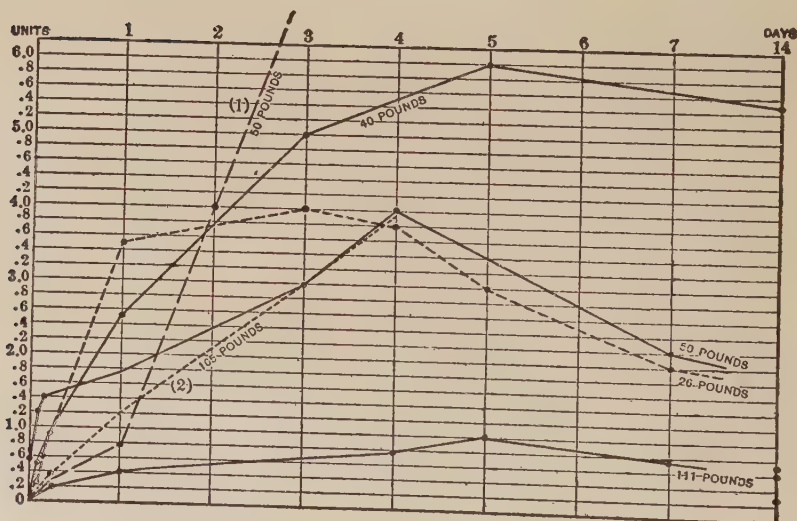
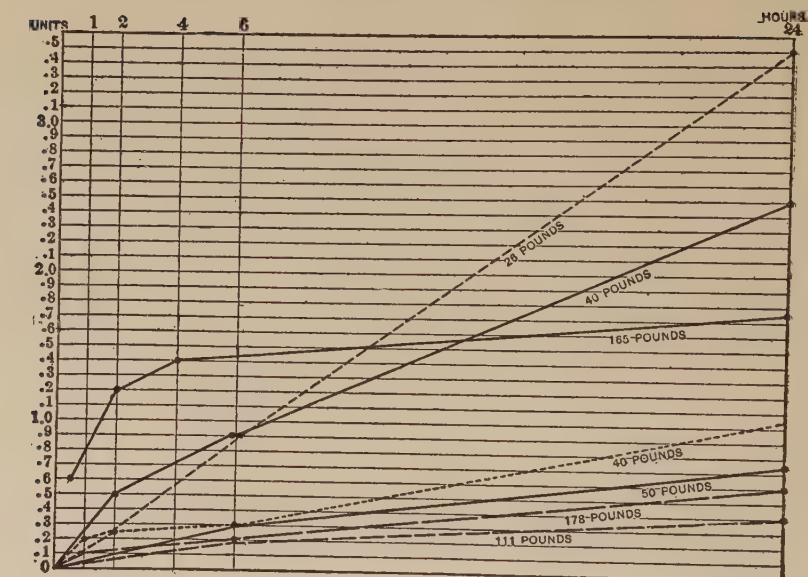


CHART I.—Showing the extent and rapidity of absorption of 10,000 units of antitoxin given subcutaneously. Each line represents the antitoxin content of 1 c. c. of blood at different intervals of time. (From Park and Biggs, *loc. cit.*)

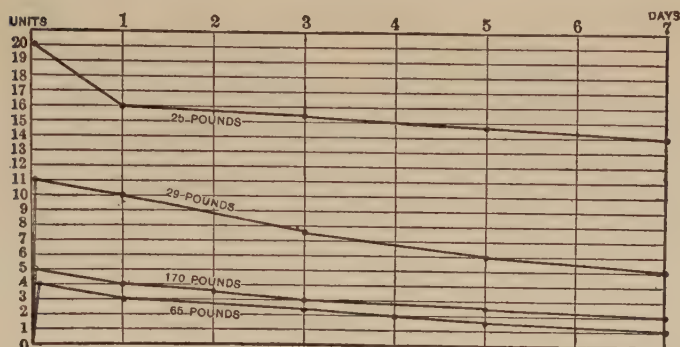


CHART II.—The antitoxic power of human blood after an intravenous injection of 10,000 antitoxic units. (From Park and Biggs, *loc. cit.*)

utes, 100 after 40 minutes, and 150 units each after 60 and 80 minutes. This rabbit died. Another animal given the same dose of toxin received 200 units of antitoxin twenty minutes later and lived. The amount necessary to save life in rabbits receiving ten fatal doses intravenously was as follows:

| | |
|-----------------------------|-----------------------|
| Given after 10 minutes..... | 5 units antitoxin |
| Given after 20 minutes..... | 200 units antitoxin |
| Given after 30 minutes..... | 2,000 units antitoxin |
| Given after 45 minutes..... | 4,000 units antitoxin |
| Given after 60 minutes..... | 5,000 units antitoxin |
| Given after 90 minutes..... | No amount |

These extremely important experiments of Park and Biggs bear out the opinion of Schick and show beyond question that the proper way to give antitoxin is to give a single adequate dose as early as possible. They emphasize the fact that probably the most important single point in the specific therapy of diphtheria is the speed with which the diagnosis can be made and the antitoxin given. At the Department of Health the dosage now employed, as given by Park and Biggs, is the following:

| | UNITS IN CASES | | | |
|----------------------------|----------------|----------|--------|-------------|
| | Mild | Moderate | Severe | Very Severe |
| Infants under 1 year..... | 2,000 | 3,000 | 10,000 | 10,000 |
| Children 1 to 5 years..... | 3,000 | 5,000 | 10,000 | 10,000 |
| Children 5 to 9 years..... | 4,000 | 5,000 | 10,000 | 15,000 |
| Persons over 10 years..... | 5,000 | 10,000 | 10,000 | 20,000 |

PRACTICAL CONSIDERATIONS CONNECTED WITH DIPHTHERIA ANTITOXIN PRODUCTION AND STANDARDIZATION

The conditions which govern the active production of toxins by bacteria in culture media are not only of great theoretical interest but possess unusual practical value in that the most important factor for the successful production of a strong antitoxin consists in the preliminary preparation of a potent toxin. The bacterial true toxins are all "exotoxins" in that they are soluble, moderately diffusible substances which pass readily from the bacterial bodies to the environment, and for this reason can be obtained most readily by the cultivation of the bacteria upon fluid media and subsequent filtration of the cultures through earth or porcelain filters.

The choice of culture or strain is an important element in this procedure, since within the same species of toxin-producing microorganisms there is much variation in the speed and energy of toxin production. Thus for unknown reasons some strains of diphtheria bacilli will far outstrip others in this respect. An excellent illustration of this is the experience of Park and Williams¹⁴ with two diphtheria cultures—a very virulent and a very weak one. Of the former, 0.002 c. c. of a forty-hour bouillon culture killed a guinea pig, while of the latter 0.1 c. c. of a similar culture was necessary for the same result.

In the case of tetanus, cultural differences do not seem to be as common. Individual strains also may gain or lose in toxin-producing powers, according to the method of handling them which is practiced. It is stated,¹⁵ for instance, that a diphtheria culture will lose in energy of toxin production if permitted to grow without sufficiently frequent transplantation. However, transplanted on solid media with reasonable frequency, these bacteria show a remarkably constant toxin production. A well-known strain, the Park-Williams No. 8, now in use in many antitoxin laboratories throughout the world, has persisted for over 15 years in producing a strong toxin. There are occasional strains among toxin-forming species which are entirely devoid of this property. Diphtheria bacilli which were virulent while possessing all the other cultural characteristics of the group have been described, but appear, from the experience of the writer, to be rather uncommon.¹⁶ Of tetanus bacilli little is known in this respect.

Given a powerfully toxic strain of the proper bacteria the method of cultivation is also of great importance in influencing the eventual yield of poison. These relations have naturally been studied with the greatest care in the case of diphtheria and tetanus

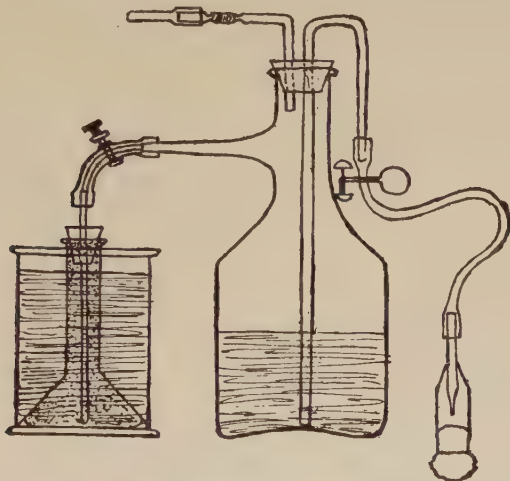
¹⁴ Park and Williams. "Pathogen. Micro-organ.," N. Y., 1910.

¹⁵ Park and Williams. *Loc. cit.*

¹⁶ Zinsser. *Jour. Med. Res.*, N. S., Vol. 12, 1907.

bacilli, since in these cases there has been the greatest practical application for such knowledge.

In the case of diphtheria, though toxin will be produced on all media on which the bacillus grows easily, the most favorable medium for this purpose is a slightly alkaline broth made of lean beef or veal infusion and containing peptone. Since acid formation hinders the production of toxin, Martin¹⁷ has suggested fermentation of the



APPARATUS ARRANGED FOR THE STERILE FILTRATION OF DIPHTHERIA CULTURES IN TOXIN PRODUCTION.

(After Rosenau, *U. S. Hyg. Lab. Bull.* 21, 1905, p. 38.)

muscle sugar with yeast, while Theobald Smith¹⁸ recommends preliminary fermentation with *Bacillus coli*.

Park and Williams¹⁹ regard this as unnecessary. They recommend a 2 per cent. peptone broth made of veal. This is neutralized to litmus and 7 to 9 c. c. of normal NaOH solution to the liter are added. In such a medium at 37.5° C. the production of toxin begins within 24 hours and reaches its highest point in from five to ten days. When at its height the process must be stopped and the cultures exposed to a lower temperature, otherwise rapid deterioration takes place because of the instability of the toxin. Even when kept cold and in the dark this deterioration proceeds steadily though slowly. At first, however, even under these conditions a comparatively extensive loss of toxin goes on—a process sometimes spoken of as “maturing of the toxin”—after which the poison strikes a

¹⁷ Martin. *Ann. Past.*, 1896.

¹⁸ Th. Smith. *Journ. Exp. Med.*, IV, 1899, p. 373.

¹⁹ Park and Williams. *Journ. Exp. Med.*, Vol. 1, 1896.

fairly constant and very gradual rate of weakening, and is, comparatively speaking, stable.

In the United States Hygienic Laboratory in Washington, according to Rosenau, the recommendations of Theobald Smith are largely followed in the production of toxin. The procedure is as follows:

The culture medium, "Smith's Bouillon," is prepared from chopped beef from which fat and tendon have been cut out. This is adjusted by phenolphthalein titration to 0.5 per cent. acidity. It is then placed into Fernbach flasks and inoculated on the surface with a Park-Williams bacillus No. 8. The flasks are incubated for 7 days at 37.5° C. The reaction of the medium after such incubation is determined, and flasks showing an acidity of 1.5 or over are discarded. The usual reaction at the end of incubation is 0.6 to 0.8 per cent. acidity. This broth is filtered through Berkefeld filters or porcelain candles.

Toxin so prepared is now tested and its L_0 and L_+ doses determined by the methods described above. Rosenau²⁰ states that poisons are discarded as containing too large a proportion of toxin if the difference between L_0 and L_+ is greater than 15 M L D. The toxin is now set aside in flasks for the process which Rosenau calls "seasoning." At intervals of about a month it is retested and finally it is found that the rate of toxoid formation decreases and the poison reaches a period of equilibrium. It can now be used for accurate determination of the L_+ dose, and this is done from careful measurements on a large number of guinea pigs.

Examples²¹ of such measurements, abbreviated for the sake of simplicity, are given in the following tables:

Toxin Determinations of M L D or "T"

| Dose in c. c. | Result |
|---------------|--------------------------------|
| 0.03 | = death in 1½ days |
| 0.02 | = death in 1½ days |
| 0.01 | = death in 2 days |
| 0.008 | = death in 3 days |
| 0.006 | = death in 3½ days |
| 0.005 | = death in 4 days M L D |
| 0.004 | = death in 6 days |
| 0.003 | = death in 8 days |
| 0.002 | = late paralysis |
| 0.001 | = well in 16 days. |

Toxin Determination of L_+ Dose

| | |
|-------------------------------|----------------------|
| 1 Antitoxin unit + 0.2 c. c. | = 0 |
| 1 Antitoxin unit + 0.21 c. c. | = 0 = L_0 |
| 1 Antitoxin unit + 0.22 c. c. | = local infiltration |

²⁰ Rosenau. *Hyg. Lab. Bull.* No. 21, April, 1905.

²¹ Examples are taken from measurements reported by Rosenau, *loc. cit.*

| |
|--|
| 1 Antitoxin unit + 0.23 c. c. = fatal in 17 days |
| 1 Antitoxin unit + 0.24 c. c. = fatal in 14 days |
| 1 Antitoxin unit + 0.26 c. c. = fatal in 9 days |
| 1 Antitoxin unit + 0.28 c. c. = fatal in 6 days |
| 1 Antitoxin unit + 0.29 c. c. = fatal in 4 days = L₊ |
| 1 Antitoxin unit + 0.3 c. c. = fatal in 3 days |

The production of antitoxin is carried out by the graded injection of antitoxin into horses. Young, healthy horses are chosen, tested for freedom from glanders, and the first injections are made either with toxin attenuated by the addition of Lugol's solution or terchlorid of iodine, or, as in the New York Health Department, the first injections consist of mixtures of toxin and antitoxin. We take our description largely from the account given by Park.²² The first injection consists of 12 c. c. of toxin (M L D 1/400 c. c.), together with 100 units of antitoxin. After the reaction from such an injection has completely subsided—after 3 to 5 days—a second injection is given of toxin without antitoxin; then 15 c. c., 45 c. c., 55 c. c., 65 c. c., 80 c. c., 95 c. c., 115 c. c., 140 c. c., etc., the intervals between injections being about three days and depending upon the reaction of the horse and the speed with which it entirely recovers from the preceding injection. In a particular case cited by Park 675 c. c. of toxin could be given by the 60th day; in this case by the 28th day the horse was yielding 225 units to the c. c.; on the 40th day, 850 units; on the 60th day, 1,000 units.

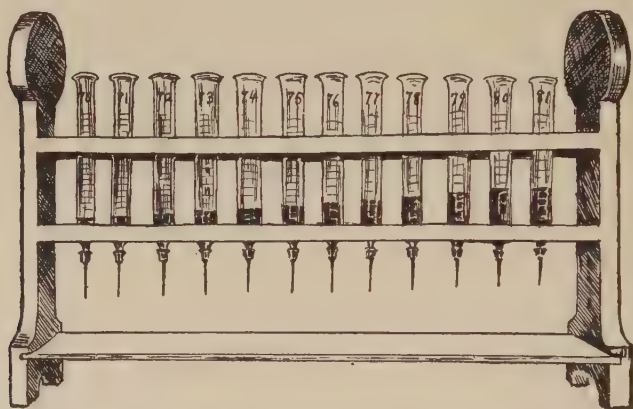
The determination of the antitoxin unit, carried out from time to time on the serum of such a horse against the L₊ dose described in our preceding table, would be carried out as follows:

In all such standardization great care must be taken in employing accurately standardized glassware. Rosenau recommends employing "capacity instruments" rather than "outflow instruments." Dilutions of unknown antitoxin are made in 0.85 per cent. sterile salt solution. As a basic dilution one part of the antitoxic serum to nine of the salt solution gives 1/10 c. c. to each cubic centimeter, and from this initial dilution further dilutions may be easily made as follows: 1 c. c. of dilution I. + 9 c. c. salt solution = 1-100, etc. A series of mixtures is then made in each of which the quantity of toxin equals the L₊ dose, and in which the quantity of antitoxin varies within a wide margin of the limits of strength to be expected. This is illustrated in the following table:

| | |
|---------------------------------------|---|
| L ₊ (0.29 c. c.) + 1/500 | c. c. of antitoxic serum = lives |
| L ₊ (0.29 c. c.) + 1/600 | c. c. of antitoxic serum = lives |
| L ₊ (0.29 c. c.) + 1/700 | c. c. of antitoxic serum = lives |
| L ₊ (0.29 c. c.) + 1/800 | c. c. of antitoxic serum = dies in 8 days |
| L ₊ (0.29 c. c.) + 1/900 | c. c. of antitoxic serum = dies in 4 days |
| L ₊ (0.29 c. c.) + 1/1,000 | c. c. of antitoxic serum = dies in 2 days |

²² Park and Williams. "Pathogenic Bacteria," p. 213.

In the above tables, according to our previous definition of the antitoxin unit, the serum would contain 900 units to the cubic centimeter, since $1/900$ c. c., injected together with the L_+ dose of the standard toxin, resulted in the death of the guinea pig in four days. In order to allow a margin of safety Rosenau and others have suggested that the unit should be determined, not by the quantity of antitoxin, which delays death by the L_+ dose for four days, but rather by the quantity which, with the L_+ dose, results in saving the life of the guinea pig. According to this latter standard the serum employed in the table would be spoken of as containing 700 units to the cubic centimeter. Of course the tabulated measurements



BATTERY OF ROSENAU SYRINGES PREPARED FOR ANTITOXIN STANDARDIZATION.
(Taken from Rosenau, *U. S. Hygienic Bulletin* 21, 1905.)

are rough, leaving an undetermined zone of 100 units. The exact number of units to the cubic centimeter could, of course, be determined with greater accuracy by now carrying out another series of tests in which the amount of serum varied between $1/700$ and $1/900$ of a cubic centimeter.

In carrying out such a standardization the toxin is diluted so that the L_+ dose is contained in 2 c. c. This can easily be done. For instance, in the above the L_+ dose being 0.29, it merely necessitates adding to each 0.29 c. c. of toxin 1.71 c. c. of salt solution, to each 2.9 c. c., 17.1 c. c. The antitoxin also is made up in such a way that the required dilution is contained in two cubic centimeters, since a total volume of 4 c. c. has been agreed upon as standard for these tests, the injected volume having much influence upon the speed of absorption. In using the so-called Rosenau syringe, shown in the figure for the standardization of antitoxin, the antitoxin is made up to 1 c. c. in each case, so that 1 c. c. of salt solution may be added to wash out the syringe after injection of the mixture. The mixtures

can be made directly in these syringes or in test tubes, and are allowed to stand one hour at room temperature, so that there may be time for complete union. If the mixtures are made directly in the syringes the needles are dipped into sterile vaselin, which closes them and prevents leakage while standing. The mixture is then forced out of the syringe with a rubber bulb, thus ensuring complete injection of all the fluid. As Rosenau states, much depends on the guinea pigs. They must be of standard weight, about 250 grammes, well fed and cared for, and must not be descendants of pigs that have shown marked or unusual resistance to diphtheria toxin. This, as Theobald Smith has shown, occasionally happens.

The antitoxic serum as obtained from the horse directly may be concentrated in a number of ways, representative of which is the method developed at the New York Department of Health by Gibson,²³ Banzhaff, and others.²⁴ The original method consisted in heating horse serum to 56° C. for 12 hours, by which some of the pseudoglobulin was converted into euglobulin, the antitoxin remaining in the pseudoglobulin fraction. After this an equal volume of saturated ammonium sulphate solution is added and the globulin precipitated. After several hours the precipitate is filtered off and again taken up in water corresponding in amount to the original volume of serum. After filtration this solution is precipitated with ammonium sulphate and this precipitate is treated with saturated solution of NaCl in quantity twice that of the original serum. After standing for 12 hours the supernatant fluid containing the antitoxin is decanted, and this is precipitated with 0.25 per cent. acetic acid. The resulting precipitate is dried by pressing it between filter papers and is placed in a parchment dialyzing bag, after neutralization with sodium carbonate. At the end of seven or more days of dialyzation against running water, the globulin solution remaining in the dialyzer is filtered and made isotonic.

More recently the method as modified by Banzhaff is as follows: The serum, as obtained from the horse, is diluted by one-half the volume of water, and to this a saturated solution of ammonium sulphate is added up to 30 per cent. saturation. This is heated to 61° C. for two hours. It is then filtered and the residue on the filter paper, which contains the antitoxin, is thoroughly dried by pressing between filter papers and is directly dialyzed.

Observations by Park and Throne²⁵ have shown that this concentrated antitoxin which, according to Gibson, represents a yield of about 70 per cent. original antitoxic power of the serum, is equally efficient for therapeutic purposes as an unconcentrated preparation

²³ Gibson. *Journ. of Biol. Chem.*, Vol. 1, 1906.

²⁴ Gibson and Collins. *Journ. of Biol. Chem.*, Vol. 3, 1907.

²⁵ Park and Throne. *Amer. Journ. of Medical Science*, Vol. 132, 1906.

and has the advantage of introducing less foreign protein into the human body. It retains its potency, according to Park and Throne, as long as does the whole serum.

ACTIVE IMMUNIZATION IN DIPHTHERIA WITH MIXTURE OF TOXIN AND ANTITOXIN

Recently Behring²⁶ has advocated the immunization of human beings with mixtures of diphtheria toxin and antitoxin. This method represents essentially *active* immunization with toxin rendered harmless by neutralization with antitoxin. The use of such mixtures had previously been studied with considerable care, in the case of the toxin of symptomatic anthrax, by Schattenfroh and Grassberger,²⁷ and the procedure had been used in the New York Department of Health for some years in the initial treatment of antitoxin horses. Theoretically considered on the basis of Ehrlich's opinions, one would be inclined to wonder at the fact that relatively neutral mixtures of toxin and antitoxin should possess any antitoxin-inciting properties. Behring explains the immunizing value of such mixtures by the reversible nature of toxin-antitoxin union in the animal body. He calls attention to the fact that our analyses of diphtheria toxin-antitoxin mixtures have been made entirely with guinea pigs as indicators. In studying such mixtures in other animals Behring has come to the conclusion that complete detoxication of the poison *in vitro* does not occur. He found, for instance, that a toxin-antitoxin mixture that was entirely innocuous for guinea pigs produced an active febrile reaction in an ass. In monkeys (*Macacus rhesus*) he finally found an animal in which he obtained evidence satisfactory to him that toxin may be powerfully active in the animal body, even if it has been previously mixed with antitoxin. If, for instance, he gave a monkey a mixture in which as much as 20 to 40 antitoxin units were mixed with one toxin unit, and repeated the injection two or three times, the animal died of subacute diphtheria toxin poisoning. The mixture ceased to be poisonous for monkeys only when the relation of antitoxin to toxin became one of 80 to 100 antitoxin units to one toxin unit. This final detoxication when sufficient amounts of antitoxin were used, it seems to us, may be taken as sufficient evidence that Behring's monkeys did not die of anaphylaxis.

We gather from Behring's writings that he attributes these differences in susceptibility to toxin-antitoxin mixtures in various animals to differences in the reversibility of the toxin-antitoxin complex in the bodies of the individual species.

²⁶ Behring. *Deutsche med. Woch.*, Vol. 39, No. 19, 1913.

²⁷ Schattenfroh and Grassberger. *Deuticke*, Wien, 1904; see also Schattenfroh, *Wien. kl. Woch.*, No. 39, Sept., 1913.

Human beings are less susceptible to such mixtures than are monkeys, but nevertheless more so than guinea pigs. It also appears that diphtheria bacillus carriers or such persons who, because of a previous infection, have antitoxin in their blood are much more susceptible to these mixtures than are others. Newborn children are less susceptible than are children from 4 to 15 years. Mixtures which are entirely neutral for the newborn may incite febrile reaction in older children. In all cases the injection of such mixtures is followed by a more or less active production of antitoxin.

The mixtures which von Behring advocates at present are so prepared that the toxin action upon guinea pigs is practically nil; in other words, the mixture is completely neutralized.

The method represents in purpose, and apparently in achievement, a safe process of actively immunizing against diphtheria. Heretofore the method of protecting human beings prophylactically against diphtheria has consisted in the injection of antitoxic serum. This, unquestionably a wise procedure, has nevertheless the disadvantage of bringing about an immunity of short duration only. Within 20 to 30 days the antitoxin injected may have completely or almost completely disappeared from the blood stream. Prophylactic immunization with the toxin-antitoxin mixtures, however, representing as it does an active immunization, is likely to be more prolonged in its effects. According to Behring a human being possessing 0.01 antitoxin unit in 1 c. c. of blood may be regarded as still moderately protected against diphtheria. According to his estimation a decline to this amount, in a person actively immunized by the mixtures (an estimation based upon curve measurements of treated cases), would take about two years. He has observed that horses that had been actively immunized by him, and subsequently used in agricultural work, retained measurable antitoxin values in their blood after five years without treatment.

Schreiber²⁸ and others state, also, that this method of active immunization with mixtures of toxin and antitoxin has the advantage of avoiding the anaphylactic dangers incident to the injection of antitoxin alone. Their opinion is probably erroneous, since it is most likely that whatever anaphylactic dangers there are result from the injection of horse serum rather than from the antitoxin contained in the injected substance. Moreover, the recent studies of Park have shown satisfactorily that the danger of anaphylaxis in the injection of antidiphtheritic sera is practically nil. Among 330,000 cases on record there were but five deaths.

The chief value of this new method of immunization is that it represents a safe technique for the prophylactic treatment of individuals exposed to the disease and possibly for the general prophylactic immunization of school children, nurses, physicians, etc. In

²⁸ Schreiber. *Deutsche med. Woch.*, Vol. 39, No. 20, 1913.

the case of children during the ages at which they are most susceptible to the disease, the prolonged immunity resulting from the treatment should strongly recommend it as a method of promise for the gradual eradication of epidemics. Behring also suggests it as a hopeful method of treatment in the case of bacillus carriers.

Schreiber and others have reported upon the effects of treatment when carried out with Behring's mixtures. In the earlier experiments of Hahn, mixtures were used in which there was a slight excess of toxin. The later experiments were made with mixtures which were completely neutralized for guinea pigs. In Schreiber's cases from two to six injections were made at intervals of three to five days, most of them subcutaneously, and some of them intramuscularly. In no case were there serious reactions, although occasionally there were slight swelling of regional lymph nodes and a little fever. The effects of immunization were noticeable about 23 to 25 days later. When two injections only had been made, at least 0.075 of an antitoxin unit to the cubic centimeter was present. The highest value obtained after two injections was one unit to one cubic centimeter. In nine patients who had been treated by four to seven injections with gradually increasing doses, as much as 10 to 75 antitoxin units to the cubic centimeter resulted. It appears, therefore, that in medical practice this method is safe, and that with as little as two injections antitoxin values may be obtained which entirely suffice for the protection of human beings against the ordinary dangers of diphtheria infection, an immunity which, as far as we can judge at present, may last about two years.

Another advantage which Behring claims for his method is the production of homologous antitoxin in human beings for the passive immunization of other human beings. Mathes has tried this in children with the idea of thereby avoiding the dangers of anaphylaxis. Incidentally it was claimed in this case that the passive immunization, when carried out with homologous serum, lasted longer than did that conferred by horse serum. However, one case is hardly enough to establish such a fact.

THE INTRACUTANEOUS METHOD OF DETERMINING TOXIN AND ANTITOXIN VALUES

Marks²⁹ was the first to utilize the prevention of local edema or injury for the determination of antitoxin values. He mixed diphtheria antitoxin and toxin and injected them subcutaneously into guinea pigs, claiming that this method was considerably more delicate than the Ehrlich method, since the amount of toxin capable of causing localized edema amounted to as little as one-twentieth of a

²⁹ Marks. *Centralbl. f. Bakt., Orig.* Vol. 36.

minimal lethal dose. This method has many points in its favor, and has been recently utilized and improved upon by Römer.

Römer^{30 31 32} has developed a method of diphtheria antitoxin standardization which depends upon intracutaneous injections into guinea pigs. The principle of this test consists in the observation that, when very slight amounts of diphtheria toxin are injected intracutaneously into the abdominal skin of guinea pigs, small areas of local necrosis result within about 48 hours. When such injections are made with mixtures of toxin and antitoxin the presence of free toxin is indicated by the appearance of such necrosis.

Before proceeding to the standardization by this method it is necessary to determine the "limes-necrosis" (just as Ehrlich determines his L+ dose), that is, the amount of toxin which, together with a given amount of toxin (1/50, 1/200, or 1/2,000), will still produce a minimal amount of necrosis after intracutaneous injection into guinea pigs. It is necessary, therefore, arbitrarily to choose a certain definite fraction of an antitoxin unit and mix this with varying amounts of toxin and inject the mixtures into guinea pigs intracutaneously. Those mixtures in which the toxin is fully neutralized will give rise to absolutely no lesion further than, possibly, a slight local edema. Those in which there is a large excess of toxin will cause extensive necrosis. Between the two, in the series, there will be a mixture in which slight local necrosis results from the injection. In this mixture the amount of toxin, just sufficient to cause noticeable necrosis in spite of admixture with the antitoxin, contains the L-n (limes necrosis) dose.

When this has been determined, then unknown antitoxin can be similarly measured against this L-n dose of the standard toxin. The method has the advantage of permitting one to work with very small quantities, since only a small fraction of a cubic centimeter need be used for intracutaneous injections; also it permits great economy of animal material, since four or five tests can be simultaneously carried out upon the abdominal wall of the same guinea pig.

The technique is not easy. We have found in studying this method in connection with some work carried on in our laboratory by Dr. M. C. Terry, that a considerable amount of practice and experience is necessary, both in carrying out the procedure accurately and in judging the lesions. However, when carefully and consistently done by an experienced worker, this method gives results which correspond with fair accuracy to measurements made of the same antitoxin by the Ehrlich method. This has been the experience of Lewin,³³ and also of Terry in the few experiments carried out by him.

³⁰ Römer. *Zeitschr. f. Imm.*, Vol. 3, p. 208, 1909.

³¹ Römer and Sames. *Ibid.*, p. 344.

³² Römer and Somogyi. *Ibid.*, p. 433.

³³ Lewin. *Centralbl. f. Bakt., Orig.* Vol. 67, 1913.

The Römer method has been recently used by clinicians for the determination of the presence of free toxin or antitoxin in the circulating blood of patients suffering or convalescent from diphtheria. Römer himself suggested this, since his method is adapted to the determination of extremely slight amounts of either substance. A recent study by Harriehausen and Wirth³⁴ illustrates the results obtained in such tests. Normal human serum injected intracutaneously into guinea pigs never caused necrosis. Neither did the similar injection of the sera of children suffering from varicella and other diseases. Of twelve children suffering from diphtheria, however, serum taken before the administration of antitoxin caused necrosis upon intracutaneous injection into guinea pigs, in every case. In spite of the administration of antitoxin, toxin was demonstrable in the blood in five cases as long as the 35th day. Of ten cases of post-diphtheritic paralysis, toxin was demonstrated in the blood of five.

Since this method of determining antitoxin values in the blood of human beings is of considerable importance and may have much practical value, it may be useful to insert an example of such an application of this method as used by Hahn³⁵ in a series of investigations mentioned elsewhere.

The standard toxin was obtained from Marburg. In a series of guinea pigs a determination was made of the smallest quantity of this standard poison which would produce just noticeable necrosis of the skin if injected into the pig intracutaneously, together with 1/2,000th of a unit of a standard antitoxin. The toxin and antitoxin were left together for 24 hours before injection, 3 hours in the incubator, and 21 hours in the refrigerator.

When this quantity of the antitoxin had been determined, it could be used in similar experiments and similarly mixed with varying amounts of the patient's serum. The amount of antitoxin present in such serum could then be easily computed. For, let us suppose that this amount of toxin, together with 1/500th of a c. c. of the serum injected intracutaneously into the guinea pig, gave the same amount of necrosis in the same time as the identical quantity of the toxin, together with 1/2,000th of a standard unit. Then 1/500th of a patient's serum was equivalent to 1/2,000th of a standard unit, and the patient's serum would contain 0.25 of a unit per cubic centimeter.

Michiels and Schick³⁶ have carried out intracutaneous reactions with diphtheria toxin directly upon the human body to determine whether or not diphtheria immunity was present. They injected 0.1 c. c. of a 1 to 1,000 dilution of toxin and claim that a positive

³⁴ Harriehausen and Wirth. *Zeitschr. f. Kinderheilkunde*, Vol. 7, 1913.

³⁵ Hahn. *Deutsche med. Woch.*, Vol. 38, No. 29, 1912.

³⁶ Michiels and Schick. *Zeitschr. f. Kinderheilkunde*, Vol. 5, 1912.

intracutaneous reaction with this amount indicates an absence of antitoxin from the blood, or at least an insufficient protection. The Schick reaction is at present carried out at the New York Department of Health, under the direction of Park, with 1/50th M L D intracutaneously injected. The dilutions are so made that this quantity is contained in a total volume of 0.1 c. c.

TETANUS ANTITOXIN AND ITS STANDARDIZATION

The methods employed in the production and standardization of tetanus toxin are in every way analogous to those used in the case of diphtheria antitoxin. A strong toxin is obtained by growing the organisms under anaërobic conditions on suitable media. According to Vaillard and Vincent³⁷ it is essential that the media upon which the tetanus bacilli are grown should be freshly made and sterilized. Apparently this precaution, which has been similarly recommended by Wladimiroff, Novy, and others, is made necessary by the gradual absorption of oxygen which takes place if the media are allowed to stand for a long time without heating. It is further necessary in preparing tetanus toxin that the culture medium should not be acid, and a weakly alkaline initial titre is advised. For the same reason, also, most workers have advised against the use of glucose or other carbohydrates in the media, since the acid formed by the fermentation of these substances inhibits growth and toxin production. Recently Hall³⁸ has advised the use of a simple meat extract broth to which have been added 1 per cent. of dextrose and 0.5 per cent. of finely powdered magnesium carbonate. The last-named substance, by neutralizing any acid that is formed from the glucose, prevents the harmful acidity. Anaërobic conditions are obtained by growing the organisms under a layer of oil in tightly stoppered flasks.

Although mice were formerly used in the standardization of tetanus toxin and antitoxin, the more recent usage has been to substitute guinea pigs as in diphtheria standardization. According to the recent directions of Rosenau and Anderson³⁹ the purposes of the standardization are carried out as follows:

The unit of antitoxin is arbitrarily designated as 10 times the smallest amount of serum necessary to preserve the life of a guinea pig weighing 350 grams for 96 hours, when given together with an official test dose of toxin. The test dose of toxin contains 100 minimal lethal doses. And the minimal lethal dose is measured against a 350-gram guinea pig.

³⁷ Vaillard and Vincent. *Ann. Past.*, 1891.

³⁸ Hall. "Univ. of Cal., Publ. in Path.," Vol. 2, No. 11, 1913.

³⁹ Rosenau and Anderson. *U. S. P. H. Service Hyg. Lab. Bull.* 43, 1908.

In carrying out the standardization the L_+ dose of the toxin is used, but, unlike diphtheria standardization, in this case the L_+ dose means an amount of toxin which will kill a guinea pig of 350 grams in four days, although united with 0.1 unit of antitoxin (it must be noted that the L_+ dose in this case is measured against one-tenth unit of antitoxin rather than against 1 unit, as in the case of diphtheria).

In determining the value of an unknown antitoxin, mixtures are made, each containing the L_+ dose of the toxin and varying quantities of antitoxin. As in diphtheria measurements, the various injection volumes are brought to 4 c. c. with salt solution, and are then injected subcutaneously into guinea pigs of about 350 grams. The table given below is taken from the Bulletin of Rosenau and Anderson.

| No. of guinea pig | Weight of guinea pig (grams) | Subcutaneous injection of a mixture of | | Time of death |
|-------------------|------------------------------|--|-------------------|-----------------|
| | | Toxin (Test dose) (gram) | Antitoxin (c. c.) | |
| 1 | 360 | 0.0006 | 0.001 | 2 days 4 hours |
| 2 | 350 | 0.0006 | 0.0015 | 4 days 1 hour |
| 3 | 350 | 0.0006 | 0.002 | Symptoms |
| 4 | 360 | 0.0006 | 0.0025 | Slight symptoms |
| 5 | 350 | 0.0006 | 0.003 | No symptoms |

In this experiment 0.0015 equals 0.10 antitoxin unit.

ANTITOXINS AGAINST SNAKE POISONS (Antivenin)

Antitoxins against snake poisons have been produced by a number of different workers, but the subject has been most extensively studied by Calmette. As early as 1887 Sewall⁴⁰ succeeded in increasing the resistance of pigeons to snake poison. Later Calmette and Physalix and Bertrand independently succeeded in producing immunity in rabbits and guinea pigs with the poison of the cobra. The serum of animals treated with snake poisons gradually acquires antitoxin properties, but the process of immunization is not a simple one, and considerable time is needed for the immunizations.

Snake poisons, as we have seen, have attracted considerable attention because of their peculiarities in being antigenic and yet differing in heat resistance and a number of other properties from the

⁴⁰ Sewall. Cited from Calmette.

bacterial toxins. It was with snake poisons that Calmette definitely showed that the union of toxin and antitoxin is a true neutralization and is not accompanied by the destruction of the toxin. These experiments, as we have seen, were elaborated later by Morgenroth, who succeeded in producing the snake poison HCl combination. It is these poisons also that have been the subject of extensive study by Flexner and Noguchi, by Kyes, and later by von Dungern and Coca. This work has been sufficiently discussed in other places and need not occupy us here. The important poisonous snakes may be divided into the colubridæ, to which class the cobra belongs, and the viperidæ, which includes the ordinary European vipers, the rattlesnake, and most of the poisonous snakes of North and South America. According to Calmette the poison of the cobra is much more heat-stable than that of the rattlesnake. Pharmacologically the poisons of these two main classes of snakes show considerable difference. In the case of the cobra there is very little local disturbance and the systemic symptoms dominate the clinical picture. Calmette describes the cobra bite as being followed only by a feeling of stiffness at the site of the bite, followed very soon by great general weakness, difficulty in respiration, slow heart action, and finally death with unconsciousness. In the case of the vipers the local symptoms are very much more marked, there being great pain and swelling and apparent clotting of the blood about the point of the bite, with a rather slower onset of systemic symptoms. In a description by Sparr⁴¹ of a case of bite by Russell's viper there was almost immediate swelling of the limb with a faint bluish tint around the pin-point puncture, and within 15 minutes great weakness, restlessness, and retching. In spite of very active local treatment, within a short time after the bite, the patient died within 24 hours of asphyxia and heart failure.

According to Calmette 0.0002 gm. of cobra poison will kill a guinea pig; Noguchi states that 0.0005 gm. of rattlesnake venom will kill a guinea pig of 250 gr. within 24-30 hours when injected intraperitoneally. The snake poisons apparently contain substances which are especially active upon nerve cells (neurotoxins), and hemolysins which act particularly upon the red blood cells. Flexner and Noguchi⁴² also speak of another poison which acts particularly upon the endothelium of the blood vessels producing hemorrhages.

According to Calmette the antisera which are produced by immunization with cobra poison are most strongly potent against neurotoxic poisons of the colubridæ and, to a certain extent, against some of the poisons of the vipers. However, the action of the cobra antitoxin against viper poison seems at best to be weak. On the other hand, antitoxins produced with rattlesnake poison are not potent against the cobra venom since, as Calmette states, the rattlesnake

⁴¹ Sparr. *Biochem. Bull.*, Dec., 1911, No. 2.

⁴² Flexner and Noguchi. *Univ. of Pa. Bull.*, Vol. 15, 1902.

poison contains hardly any neurotoxin. Antitoxins may be produced by the gradual immunization of horses, and have been produced in this way by Calmette in the Pasteur Institute of Lille for some years. Calmette standardizes his antitoxin by determining the amount of serum which completely neutralizes *in vitro* 0.0001 gm. of the poison as tested upon white light. He also determines the protective power by injecting a rabbit with 2 c. c. of the serum and two hours later gives 1 gm. of the poison.

Noguchi has studied rattlesnake poison particularly and succeeded in preparing a strong antitoxin by the gradual immunization of a goat. Great difficulty has always been experienced in attempts at immunization with rattlesnake poison because of the very violent local injury produced by injections of the venom. The potency of the serum produced by him was such that 2½ c. c. of goat serum protected guinea pigs against 12 times the fatal dose of rattlesnake poison if given at the same time. If the antivenin was given one hour later, 5 times the amount of serum had to be given.

PASSIVE IMMUNIZATION IN DISEASES CAUSED BY BACTERIA WHICH DO NOT FORM SOLUBLE TOXINS

As we have stated in another place the greatest therapeutic successes with passive immunization have been achieved in bacterial diseases in which the malady is essentially a toxemia due to a soluble toxin. In such cases the serum of actively immunized animals contains specific antitoxins by virtue of which the toxins circulating in the blood of the patient are directly neutralized, quantity for quantity, with consequent therapeutic benefit. In the case of bacteria in which no toxins are formed, the immunization of an animal is not followed by the formation of any poison-neutralizing principle. Here the injection of bacteria, dead or alive, or the invasion of the bacteria in the course of spontaneous disease, is followed by the formation of specific antibacterial substances, lytic, opsonic, agglutinating, or precipitating bodies, the nature of which we have discussed in other chapters. The toxemia which occurs in such cases is due as we have seen to derivatives of the bacterial protein which by some observers are regarded as preformed endocellular poisons liberated by the lytic action of the serum, and by others as split products of the bacterial protein, non-existent until the bacterial cell has been acted upon by the serum components and destroyed. However this may be, the recovery from diseases of this nature is accomplished by bacterial destruction; this may be direct, by the bactericidal action of the serum, or indirect by opsonic properties which induce phagocytosis. The poisons which are liberated from the bacterial bodies, if free, can do their injury, and no neutralizing sub-

stance is formed in the body fluids to prevent their action as far as we know. Immunity in such cases, then, is not an antitoxic immunity in any sense of the word; it is rather an antibacterial immunity in which the disease is prevented or cured only when complete destruction of the bacteria has taken place. If an animal or a human being is prophylactically immunized against diseases of this kind (typhoid, cholera, etc.), it is easy to see that an increased presence of antibacterial substances, bactericidal or opsonic, in the circulation would serve efficiently and rapidly in disposing of the small numbers of invading micro-organisms which ordinarily enter the body in spontaneous infections. And, indeed, experience has shown that prophylactic immunization can be successfully carried out in the case of cholera, typhoid fever, plague, and other diseases which are sufficiently prevalent endemically or epidemically to justify prophylaxis on an extensive scale.

However, when in diseases of this kind the body is already extensively infected and has begun, as is usually the case, to respond spontaneously with the formation of specific antibodies, it has been a matter of doubt whether or not passive immunization, that is, the introduction of specific antibodies in the form of the serum of a highly immunized animal, is therapeutically of value. Indeed, it has been feared that the use of such sera may even be harmful in that the sudden introduction of large amounts of bactericidal substances might lead to a sudden liberation of large quantities of poisonous products and consequent rapid toxemia.

The conditions in such cases are exceedingly complex and many gaps exist in our knowledge concerning them. The bacteria when invading the body, immediately enter into conflict with the protective forces, as we have stated in the chapter on Infection. If a considerable degree of resistance exists, let us say as the result of preceding immunization or a recent attack of the disease, there is a rapid destruction of the bacteria, probably by active phagocytosis. It has been shown by Bordet in the case of cholera and more recently by Gay with typhoid, that injection of the organisms into immunized animals is followed by prompt and high leukocytosis, whereas similar injections into normal animals usually induce a temporary leukopenia. When the invaded animal is not particularly resistant the bacteria may accumulate and, as in the case of pneumococci and streptococci, develop phagocytosis-resisting properties (capsule formation, etc.); or, as in the case of typhoid bacilli, there may be an immediate liberation of toxic substances (anaphylatoxins) by reaction between bacterial cell and blood plasma, which can induce leukopenia, and by this means the organisms may be protected from phagocytic destruction. Experience with curative sera in all of the conditions of this class has yielded promising results only when the cases have been treated with the sera at early stages of the disease,

either when the invading germ was still localized or, at least, when the septicemic condition was not yet thoroughly established. It may be that the doses heretofore given have been insufficient, and indeed recent experiences with pneumonia seem to indicate that this may have been, in part, the cause of earlier failures. Yet in pneumonia the septicemia probably does not represent the firm establishment of a foothold by the pneumococcus in the circulation but rather a continuous discharge of new organisms into the blood from the localized lesion in the lung.

It is our own opinion moreover that septicemia as usually observed clinically represents in most cases exactly this condition, that is, a more or less continuous discharge of the bacteria into the blood from some active focus with a continuous destruction of the organisms after they have entered the blood stream. It is only when the resistance of the body is overwhelmed, in the later stages of the disease, that the bacteria can continue to grow and develop in the circulation, and this stage probably does not occur until death is imminent. In such septicemic diseases as streptococcus infection, typhoid fever, plague, anthrax, and many others the presence of the bacteria in the blood at the time when the patient is still in a condition of powerful resistance probably means that the bacteria are being supplied to the blood from the local lesions. There is probably just such a continuous discharge of bacteria from the focus into the blood with active destruction after the bacteria have entered the circulation. This seems especially probable from the fact that in many of these diseases the protective antibodies, bactericidal and opsonic, can often be demonstrated in the blood serum in quantities higher than normal at the very time when blood culture yields positive results. In typhoid fever, of course, it is well known that bactericidal titres of over 1-50,000 are often present while the patient may still be very sick, and in the more chronic streptococcus conditions with malignant endocarditis we have often seen that opsonic properties on the part of the patient's serum against the very organism invading him are considerably higher than normal. We take this to mean that the injection of immune sera would simply aid in more rapidly freeing the blood stream of the bacteria, the cure of the disease, however, involving a destruction of the focus. This, of course, is not possible merely by the injection of the serum. When, as in some cases of streptococcus infection, the focus can be surgically reached, the septicemia will often disappear and cure result, as we have ourselves had the opportunity to observe. When the focus cannot be reached surgically, it may nevertheless be a wise procedure to inject considerable amounts of immune serum, for, by keeping the blood stream free of bacteria, the case may be influenced favorably. Pneumonia is an example of this. Former failures have recently been turned into partial success by the work of Neufeld and of Cole

merely by the use of larger quantities of immune sera essentially similar to sera used at previous times, and Cole attributes the apparently favorable results to the fact that the blood stream can be cleared of bacteria although the focus cannot itself be affected.

Cure of such diseases, therefore, by serum treatment can hardly be expected. Favorable influence of the disease by energetic serum treatment may, however, be hoped for.

In discussing this subject it must not be forgotten, however, that in most of the diseases which we have classified, on the basis of prevailing opinions, as caused by bacteria that do not form true toxins, the formation of such poisons has been claimed by a number of careful and eminent observers. In the case of the typhoid bacillus, especially, Chantemesse, Kraus and Stenitzer, and others have claimed the existence of a true toxin and a consequent antitoxin in immune sera. Similar claims have been made for the cholera spirillum by Kraus and Doerr, for the streptococcus by Marmorek, and for the plague bacillus by Markl and Rowland. Since these claims have been made on the basis of extensive experimentation by competent men the question must be left open, and the possibility of antitoxic properties on the part of the sera cannot be completely ignored. Since in most cases, however, the poison-neutralizing properties of the immune sera in this disease have not exceeded more than 1 to 2 multiples of the M L D of the bacterial poisons, it does not seem impossible that the apparent antitoxic properties may have represented merely an acquired tolerance to anaphylatoxic poisons of which we have spoken in another place.

SERUM TREATMENT IN EPIDEMIC CEREBROSPINAL MENINGITIS

Serious attempts to produce curative sera against the epidemic form of cerebrospinal meningitis were not made until 1906 and 1907, when this disease appeared epidemically chiefly in Europe, where it appeared most severely in Eastern Germany, and in the Eastern United States.

In 1906 Kolle and Wassermann immunized three horses with meningococci, using for immunization purposes the dead organisms followed by living cultures and cultures shaken up in distilled water, the so-called artificial aggressins of Wassermann and Citron. They obtained sera of considerable potency when measured against meningococcus cultures, and suggested standardizing the sera by complement fixation. They did not at this time treat human beings, but suggested the use of the serum subcutaneously and intravenously in meningitis cases. Very soon after the publication of the work of Kolle and Wassermann Jochmann⁴³ also produced an antimeningo-

⁴³ Jochmann. *Deutsche med. Woch.*, 1906, Vol. 32, p. 788.

coccus serum by immunizing horses with proved meningococcus cultures, in his cases making a polyvalent serum by the use of many different strains of the organism. The sera which he obtained were highly agglutinating, somewhat bactericidal, and, according to him, not antitoxic. He first succeeded in immunizing guinea pigs against meningococci by injecting the serum 20 hours before infecting the animals. He also treated 40 cases of meningitis in man and obtained encouraging results in cases treated before the development of hydrocephalus. Believing that possibly intraspinal injection of the serum might offer advantages, he first determined by experiments upon the dead body that the injection of methylene-blue intraspinaly passed from the point of injection in the lumbar regions as far up as the olfactory nerves. After having determined this he treated 17 cases by tapping the spinal canal, taking out 30 to 50 c. c. of spinal fluid and then injecting about 20 c. c. of the serum. Of these 17 cases only 5 died, and Jochmann expresses himself optimistically in consequence.

Meanwhile Flexner⁴⁴ had been working upon the same subject, laying a rather more thorough basis for therapy in careful animal experimentation. He produced the typical disease in monkeys by intraspinal inoculation of the meningococci and then saved the animals from death by following the infection with the injection of serum intraspinaly six hours later. In his earlier articles he expresses himself with much conservatism, but his studies were continued and extensive opportunity for testing the serum which he then produced, together with Jobling,⁴⁵ was offered by the continuance of the epidemic throughout the United States.

The results with the serum produced at the Rockefeller Institute have since then proved to be uniformly favorable. The method of intraspinal inoculation of the serum after the removal of some of the spinal fluid was the method finally adopted by Flexner as most favorable, and this is the method in current use to-day. In 1908 Flexner and Jobling reported upon 47 cases treated with the antiserum of which 34 recovered. Of 12 additional cases reported in an addendum only 4 died. In the most recent summary by Flexner⁴⁶ records of 1,294 cases that have been treated with the serum prepared at the Rockefeller Institute are analyzed. Of this number, unselected and treated in many different parts of the world, 69.1 per cent. recovered. It is of course very difficult to obtain exact comparative data on the efficiency of any method of treatment in a disease as irregular in its clinical manifestations as epidemic meningitis, especially since the mortality attending upon different

⁴⁴ Flexner. *J. Exp. Med.*, Vol. 9, 1907, and *J. A. M. A.*, 1906, Vol. 47, p. 560.

⁴⁵ Flexner and Jobling. *J. Exp. Med.*, Vol. 10, 1908.

⁴⁶ Flexner. *J. Exp. Med.*, Vol. 17, 1913.

epidemics is subject to great variations. For this reason we can draw conclusions only from a large statistical material. However, we know that the average mortality of epidemic meningitis before the introduction of specific therapy ranged certainly higher than 65 per cent., and in carefully studied epidemics usually between 70 and 80 per cent. The statistics of Flexner showing a mortality hardly exceeding 30 per cent. in unselected cases unquestionably marks a wonderful therapeutic triumph. It must be remembered in considering the benefits of this serum that in unselected cases there must be many in whom the disease has produced marked anatomical changes in the central nervous system before the serum is used. It is well known, of course, that the later manifestations of this disease, which often lead to death with hydrocephalus, asthenia, and malnutrition, are the remote results of the anatomical injuries produced by the inflammatory reactions accompanying the earlier manifestations of the acute infection. These conditions of course cannot be expected to yield to serum treatment. It must be assumed, therefore, that were we able to obtain statistics of cases diagnosed and treated soon after the onset the figures would be even more favorable than those stated above.

The action of the serum seems very largely to be an opsonic one, in that, under the influences of serum, a powerful phagocytosis of the meningococci takes place. It is also possible that to a certain extent bactericidal action participates, in that the injection of the serum into the closed space may give rise to a sort of intraspinal Pfeiffer reaction with energetic ingestion of the bacteria by leucocytes.

The standardization of the antimeningococcus serum has been worked out particularly by Jobling.⁴⁷ After attempting to standardize the sera by their protective power against meningococcus infection in animals and by complement fixation, as suggested by Kolle and Wassermann, Jobling has come to the conclusion that neither of these methods is sufficiently regular, and that the most suitable procedure is a standardization by opsonin determination. The method as worked out by him depends upon determining the highest dilution of the immune serum at which opsonic action against the meningococcus is still evident. He suggests as a definite and suitable standard of strength opsonic activity at a dilution of 1-5,000 of the antiserum.

SERUM TREATMENT IN STREPTOCOCCUS INFECTIONS

The attempts to produce powerful immune sera against streptococci date back to the earliest days of immunology. That the subject is a particularly difficult one follows from the great confusion

⁴⁷ Jobling. *J. Exp. Med.*, Vol. 11, 1909.

which has prevailed, and, to a great extent, still prevails, regarding the classification of the streptococci and their interrelationship. There are apparently a large number of different strains of streptococci which vary from each other, not only culturally, but also in regard to agglutination and bactericidal reactions. For this reason it is not at all a foregone conclusion that a serum prepared by the immunization of an animal with a streptococcus of one type will have any protective action against other strains. The subject has been still more complicated recently by the discovery of Rosenow⁴⁸ that the various types of streptococci (viridans, hemolyticus, etc.) are not constant in their properties, but may be artificially transformed one into the other, and that even mutation of true pneumococci into true streptococci may take place. Most important in this connection is the observation that a pneumococcus sent to Rosenow was altered by him by special methods of cultivation in such a way, that not only its morphological and cultural properties were changed, but also its agglutination reactions. These observations are of the utmost importance in connection with attempts at producing specific sera which can be utilized therapeutically. In all cases, therefore, in which streptococcus immune serum is at all used it must be remembered that the disease produced in human beings by organisms classified among the streptococci are by no means necessarily closely related in biological reactions, and the same immune serum may be extremely potent in one case and entirely useless in another.

That animals could be successfully immunized against streptococci was shown early in the history of investigations in immunity by a number of workers, notably Roger, Behring, von Lingelsheim, and Mironoff. The first extensive attempts to produce a curative serum for use in passively immunizing human beings were made by Marmorek⁴⁹ at the Pasteur Institute in 1895. The basic idea from which Marmorek worked was the similarity of all the streptococci producing disease in human beings. He also believed that the most powerful serum could be produced with cultures whose virulence had been greatly enhanced by animal passages. When such cultures were grown on mixtures of human serum and broth he asserted furthermore that soluble poisons were produced which could be obtained by filtration of the culture fluids. For these reasons he immunized horses with cultures rendered highly virulent by very gradual injections first of dead then of living organisms, finally injecting also considerable quantities of culture filtrates.

Testing these sera upon animals, he was successful in protecting against streptococcus infection when the serum was administered 12 to 18 hours before the bacteria were injected. He expressed the opinion that the serum was antitoxic as well as antibacterial. In

⁴⁸ Rosenow. *Journ. A. M. A.*, Feb., 1914.

⁴⁹ Marmorek. *Ann. Past.*, Vol. 9, 1895.

his earliest reports the results of the treatment of 413 cases of erysipelas leave one very much in doubt as to the value of the serum since the difference in mortality between the treated and the untreated cases is less than 2 per cent. However, an analysis of the individual cases makes the serum treatment appear more favorable. He reported good results also in 7 cases of puerperal septicemia and in scarlatinal angina. Later observers, notably Lenhartz,⁵⁰ Baginsky,⁵¹ and many others, have not been able to confirm the favorable results reported by Marmorek, and it may be stated that at the present day the value of Marmorek's serum is very much in question. Antistreptococcus sera have also been produced by Aronson⁵² and Tavel, Van de Velde, Menzer,⁵³ Moser,⁵⁴ and some others. Aronson at first worked from the idea which Marmorek also had used that there was a close relationship between the various streptococci pathogenic for man. He adopted the opinion first developed by Denys⁵⁵ and Van de Velde that many different strains should be used for immunization in order to allow for possible difference in the characteristics of the pathogenic streptococci. This principle of the necessity for the production of polyvalent sera was also emphasized strongly by Tavel, who based his opinion on careful agglutination tests, and by Menzer and Moser.

That the action of the antistreptococcus sera, however produced, is very largely due to its opsonic properties has been shown by Bordet,⁵⁶ by Meier and Michaelis, and a number of other workers. If there is any bactericidal power it is probably relatively slight.

It would be quite impossible to attempt in this place to analyze the large number of streptococcus infections of man which have been treated with one or the other antistreptococcus sera. Those mentioned, moreover, do not by any means include all the sera which have been produced and marketed for this purpose. In general we may say that here, as well as in the cases of other sera in which no antitoxic action is evident, beneficial results have been obtained chiefly in cases in which the streptococcus infection has been localized and treated early after its inception. In generalized or advanced cases it cannot be said that the results are encouraging. Even in animals, in which experimental conditions can be so much more thoroughly controlled, the protective action of even the strongest sera is evident only if the serum is administered either before in-

⁵⁰ Lenhartz. "Die Septischen Erkrankungen Hölder," Wien, 1903.

⁵¹ Baginsky. *Berl. kl. Woch.*, 1896, p. 340.

⁵² Aronson. *Berl. kl. Woch.*, Vol. 39, 1902; *Deutsche med. Woch.*, 29, 1903.

⁵³ Menzer. *Berl. kl. Woch.*, 1902, and *Münch. med. Woch.*, 1903.

⁵⁴ Moser. *Wien. kl. Woch.*, 1902.

⁵⁵ Denys. *Bull. de l'Acad. Belge*, 1896, cited from Schwoner K. and L. H., Vol. 2.

⁵⁶ Bordet. *Ann. de l'Inst. Past.*, 1897.

fection or within a very definite period after inoculation. The standardization of streptococcus sera may be accomplished by determining its protective value for animals when injected 18 to 20 hours before infection. When the sera are produced by immunization with streptococci obtained from the human body and without pathogenicity for animals the standardization is of course unsatisfactory.

SERUM TREATMENT IN PNEUMONIA

Attempts to work out a therapeutically valid method of passive immunization in pneumonia have been many and date from the very beginning of the discovery that pneumonia was a bacterial infection. Sera have even been marketed and used, but until recently no very encouraging results were obtained. Recent studies have revealed that in pneumonia the serum of convalescents contains practically no bactericidal properties for the pneumococcus, and that the protective powers of such serum depend upon the presence of immune opsonins or bacteriotropins, by means of which the pneumococci are rendered amenable to phagocytosis. Virulent pneumococci are not as a rule phagocytatable in the presence of normal serum. However, in the presence of immune serum powerful phagocytic action can be observed.

Neufeld has studied the conditions of pneumococcus immunity most thoroughly in recent years. The most important advance from a practical point of view was a discovery made by him, with Händel,⁵⁷ in 1909. They determined that there was a definite difference between various pneumococci in their reactions to immune serum; in other words, pneumococci could be grouped into various serological types. The serum produced with organisms of one type did not protect against infection with other strains. In consequence they called attention to the importance of determining the type of pneumococcus which causes the individual pneumonia so that the corresponding immune serum might be used. They produced a highly potent anti-pneumococcus serum by the injection of horses and donkeys with highly virulent pneumococci grown on fluid cultures, then determined the high protective power of this serum upon animals and used it in the treatment of patients by intravenous injection. Their results were exceedingly encouraging. In reporting their results Neufeld and Händel state that considerable doses must be given. They call attention to the fact, revealed by their animal experiments, that moderate amounts do not, as in the case of diphtheria serum, exert a correspondingly slight amount of beneficial action, but that in the case of the pneumonia serum amounts smaller than a certain

⁵⁷ Neufeld and Händel. *Zeitschr. f. Imm.*, Vol. 3, 1909, and *Arb. aus dem kais. Gesundh. Amt.*, Vol. 34, 1910.

active minimum seem to exert absolutely no beneficial action. This is a fact which later was also determined by Dochez.

In confirmation of the work of Neufeld and Händel⁵⁸ Dochez and Gillespie⁵⁹ have also been able to determine that there are at least two distinctive groups of pneumococci which differ from each other as far as agglutination and serum protection experiments are concerned. In addition to these two fixed types they separate as a third group the streptococcus or *Pneumococcus mucosus* and a fourth heterogeneous group which seems to fit in with none of the others as far as serum reactions can determine.

Cole,⁶⁰ therefore, adopts the reasoning of Neufeld in that he advises the determination of the type of pneumococcus present in cases of pneumonia as a guide to the type of antiserum to be used. The type of organism is determined as soon as a case comes under observation and 50 to 100 c. c. of the homologous antiserum is injected intravenously. The result in the few cases so far treated by Cole and his associates has been encouraging.

Protective substances, according to Dochez, appear in the serum of treated cases very shortly after the administration of the serum, whereas in untreated lobar pneumonia such protective substances usually do not appear until after the crisis. Apparently Cole believes that the great value of passive immunization of this kind in pneumonia lies in the fact that the bacteriemia shown to prevail in probably all cases of lobar pneumonia is either cured or improved by the treatment, converting the disease, which is by nature, at least for a time, a septicemia, into a localized pulmonary infection. Experience with antipneumococcus serum so far has been too limited to warrant final judgment as to its permanent place among therapeutic agencies.

THE SERUM TREATMENT OF TYPHOID FEVER

The first extensive attempts to treat typhoid fever by passive immunization with the serum of treated animals were made by Chantemesse, who immunized horses with filtrates of typhoid cultures subcutaneously, and with emulsions of virulent bacilli intravenously. Chantemesse believed that the serum of horses which had been treated in this way for very long periods possessed, not only bactericidal action, but stimulated phagocytosis, and possessed a certain limited amount of neutralizing power against the toxic properties of the typhoid filtrates. At the International Congress of Hygiene in Berlin in 1907 Chantemesse⁶¹ reported upon a thousand cases

⁵⁸ Neufeld and Händel. *Arb. aus dem kais. Gesundh. Amt.*, Vol. 34, 1910.

⁵⁹ Dochez and Gillespie. *Jour. A. M. A.*, Vol. 61, p. 727, 1913.

⁶⁰ Cole. *Jour. A. M. A.*, Vol. 61, p. 663, 1913.

⁶¹ Chamtemesse. International Congress of Hygiene, Berl., Sept., 1907; *Ref. Bull. de l'Inst. Pasteur*, Vol. 5, 1907, p. 931.

treated with his serum. Of this number 43 only died, whereas the average mortality during the same six years at the Paris hospitals was 17 per cent. The injection of the serum he claimed very markedly improved the condition of patients in that, after a preliminary period of no apparent change lasting from several hours to 5 or 6 days, the temperature goes down and the general condition of the patient changes considerably for the better. He noticed very few complications in these cases, and intestinal hemorrhage occurred four times only.

A remarkable feature of Chantemesse's treatment is that he injected into the patients a few drops only of the serum, and rarely made a second injection, facts which alone tend to persuade one that his apparent therapeutic success was a fortunate accident.

The opinion originally expressed by Chantemesse that the serum of horses vigorously treated with typhoid bacilli possesses in addition to its bactericidal and opsonic powers definite antitoxic properties recurs again in the work of a number of investigators. Besredka⁶² prepared a serum by the intravenous injection of typhoid cultures heated to 60° C., continuing the immunization for 6 months. He claims that this serum possesses what he designates as "anti-endotoxic" properties. A dry extract of typhoid bacilli which in dose of 0.01 gram killed a guinea pig of 300 grams regularly became innocuous when mixed with small quantities of this horse serum. One c. c. of the horse serum neutralized often as much as two fatal doses of the serum, but it is important theoretically to recognize that Besredka states particularly that even an increase of the quantity of serum never neutralized more than two fatal doses. This is particularly important in connection with the more recent studies on toxic split proteins by Vaughan, and on anaphylatoxins by Bessau and by Zinsser and Dwyer, in which it has been shown that an animal acquires a tolerance against the toxic substances produced from bacterial and other proteins which, however, never exceeds one or two multiples of the minimum lethal dose. This fact alone would militate against considering the serum of Besredka in any way antitoxic in the sense in which the word is used concerning diphtheria and tetanus antitoxins where neutralization of poison follows roughly the law of multiples. Besredka's anti-endotoxic sera has recently been very thoroughly investigated by Pfeiffer and Bessau.⁶³ These investigators have found that Besredka's serum exerted a very definite beneficial influence upon typhoid infection in guinea pigs if injected at the same time with the bacilli. In their experiments it also protected somewhat against the toxic properties of substances derived from the typhoid bacillus, and Pfeiffer and Bessau did not believe that this was due to a true antitoxic action, nor that the serum was

⁶² Besredka. *Ann. de l'Inst. Pasteur*, 19, 1905, and 20, 1906.

⁶³ Pfeiffer and Bessau. *Centralbl. f. Bakt.*, Vol. 56, 1910.

superior in this respect to the ordinary bactericidal sera prepared by inoculating animals with typhoid bacilli. Kraus and Stenitzer⁶⁴ have also taken up the study of typhoid immunization from the point of view that the typhoid bacillus produces a true toxin, and that therefore a true antitoxic action could be expected from the sera produced by immunization with typhoid filtrates. It should be noted that, in spite of the most common opinions against this at present, a similar point of view was advanced by MacFadyen,⁶⁵ and more recently by Arima.⁶⁶ Kraus and Stenitzer⁶⁷ immunized horses and goats very highly with extracts of agar cultures and with broth filtrates by intravenous injection. The serum which they produced in this way not only possessed the ordinary bactericidal action, but, they claimed, neutralized also toxic broth filtrates, not only of the typhoid, but of the paratyphoid bacilli. The serum of Kraus and Stenitzer has been used by a number of observers, among whom are Herz,⁶⁸ Forssmann, Unger, Russ, and others, and the results are said to be encouraging in early cases.

Rodet and Lagrifoul⁶⁹ immunized horses with living typhoid cultures, and also claim favorable results.

Mathes,⁷⁰ continuing the work of Gottstein after the death of the latter, employed the method of immunizing with the product obtained by digesting typhoid bacilli with trypsin. The poison so produced he speaks of as "fermotoxin." Lüdke⁷¹ slightly modified the Gottstein-Mathes method by digesting the typhoid bacilli with pepsin and hydrochloric acid, and with the poison so produced immunized 8 goats, reënforcing the immunization by the subsequent injection of the bacilli themselves. He claims that 0.05 to 0.1 c. c. of the serum so produced protected animals against five times the lethal dose of the poison. In a small series of human cases treated by this method he reports good results.

Garbat and Meyer⁷² immunized animals with sensitized typhoid bacilli, and claim that the most potent sera for typhoid immunization can be obtained by the combination of sera produced by the injection of sensitized and of unsensitized bacteria. They assert that the typhoid bacillus contains two definite antigens, one particularly as-

⁶⁴ Kraus and Stenitzer. *Wien. kl. Woch.*, Vol. 20, 1907, pp. 344 and 753, and Vol. 21, 1908, p. 645.

⁶⁵ MacFadyen. Cited from Stenitzer in "Kraus und Levaditi Handbuch," Vol. 2.

⁶⁶ Arima. *Centralbl. f. Bakt.*, 65, 1912, p. 183. Orig.

⁶⁷ Kraus and Stenitzer. *Wien. kl. Woch.*, Vol. 22, 1909, p. 1395; *Deutsche med. Woch.*, March, 1911.

⁶⁸ Herz. *Wien. kl. Woch.*, Vol. 22, 1909, p. 1746.

⁶⁹ Rodet and Lagrifoul. *C. R. de la Soc. de Biol.*, April, 1910.

⁷⁰ Mathes. *D. Archiv f. kl. Med.*, Vol. 95, 1909.

⁷¹ Lüdke. *D. Archiv f. kl. Med.*, 98, 1910.

⁷² Garbat and Meyer. *Zeitschr. f. exp. Path. u. Ther.*, Vol. 8, 1911.

sociated with the bacterial ectoplasm, which becomes active when the bacteria enter the animal body, and a truly endocellular poison which does not become active until the surrounding ectoplasm is dissolved. They believe that sensitizing bacteria is a method for the production of endotoxin, and think that therefore the ideal serum for the treatment of typhoid consists of a mixture of two sera produced each with one of the antigens, that is, with sensitized and unsensitized bacteria. Rommel and Herman⁷³ did not obtain encouraging results with this serum.

From a study of the literature it seems to us that in spite of the many different methods of production employed by various observers in their studies on typhoid sera it is quite likely that all these sera are essentially alike, containing, quantitatively, according to the degree of immunization, bactericidal, agglutinating, and opsonic properties, with possibly a limited amount of neutralizing power for the poisons liberated from the typhoid bacilli in the body. As far as we can judge from clinical reports the therapeutic value of the sera so far produced is not very great. It seems that cases treated early in the disease may be benefited, and possibly an early cessation of the bacteriemia can in this way be attained. However, it does not seem either theoretically or from the study of clinical publications that any very marked effects have followed the use of any of the sera in advanced cases.

THE SERUM TREATMENT OF PLAGUE

That the serum of animals immunized with killed plague cultures may actively protect normal animals from experimental infection was first shown by Yersin, Calmette, and Borrel.⁷⁴ The serum which they produced possessed apparently powerful bactericidal action, but no antitoxic properties were demonstrated. They determined its protective powers by injecting measured quantities into mice and infecting them with fatal doses of virulent plague bacilli 24 hours later. The Yersin serum which was produced for the treatment of plague as a result of these experiments was made, then, by the gradual immunization of horses with first dead plague bacilli, finally with virulent living organisms. The serum has been extensively used by many observers with results that leave one much in doubt as to its efficacy. Yersin⁷⁵ himself, reporting on an epidemic in Nhatrang, reports a general mortality of 73 per cent. for the whole epidemic, a mortality of 100 per cent. in untreated cases, and of 42 per cent. among those treated with his serum. Good results were also reported from the epidemics in Amoy and Canton in 1896.

⁷³ Rommel and Herman. *Centralbl. f. Bakt.* Ref. Vol. 53, 1912.

⁷⁴ Yersin, Calmette, and Borrel. *Ann. de l'Inst. Past.*, 1895.

⁷⁵ Yersin. *Ann. de l'Inst. Past.*, 1899.

However, these results apparently were not accepted by all observers as proving the efficiency of the serum, since the number of cases observed were few, and the irregularity in the gravity of the disease in different individuals makes statistical evidence unreliable unless large material can be studied. Kolle and Martini⁷⁶ announce that Dr. Choksy reported very poor success with the Yersin serum, and cite a number of later writers whose results with this serum were also unsatisfactory when used on human beings. That the serum unquestionably contains antibodies against the plague bacillus is testified to, not only by the French observers themselves, but also by the German Plague Commission of 1899, and by Kolle and Martini⁷⁷ themselves. The Commission experimented with this serum upon monkeys, and showed that it possessed unquestionable protective powers in rodents and in monkeys when given 24 hours before the plague infection, and in monkeys possessed fair curative properties when injected 24 hours later than inoculation with the plague bacilli. Because of the doubtful success in the treatment of human beings with this serum Yersin and Roux at the Pasteur Institute later altered their methods of serum production by injecting, not only dead and living plague cultures, but considerable quantities of culture filtrates after the horses had attained a high degree of immunity. Later observations on the Yersin⁷⁸ serum have been published by the British Plague Commission in 1908 and 1911. In this investigation the cases were controlled as to their severity by blood culture, since it had been claimed by a number of earlier investigators that the Yersin serum was efficient in mild cases, but failed entirely in the severe ones. It seems from the report of this Commission that ordinarily 70 per cent. of cases of plague without bacilli in the blood survive while three-quarters of those with mild septicemia die, and all of those with a marked septicemia succumb. In the summary given of 146 cases treated with Yersin's serum by the British Commission 65.1 per cent. died, whereas of 146 untreated controls 71.90 per cent. died. These figures, together with an analysis of the percentages, classified according to the severity of the infections, do not show a very marked curative action on the part of the serum.

Markl,⁷⁹ who claims that the plague bacillus produces a soluble toxin, has produced a plague serum by immunization of animals by filtrates of broth cultures. He claims that 0.1 c. c. of his serum, as produced at Vienna, will protect various animals against lethal doses of plague bacilli if given at the same time. He attributes much of

⁷⁶ Kolle and Martini. *Deutsche med. Woch.*, 1902, p. 29.

⁷⁷ German Plague Commission. *Arb. a. d. kais. Amt.*, Vol. 16, 1899.

⁷⁸ British Plague Commission. *Journ. of Hyg.*, Vol. 12, Sup., 1912, p. 326.

⁷⁹ Markl. *Centralbl. f. Bakt.*, 24, 1898; *Zeitschr. f. Hyg.*, 37, 1901; *Zeitschr. f. Hyg.*, 42, 1903.

its curative action to the fact that in the presence of this serum active phagocytosis takes place.

Dean,⁸⁰ in 1906, also claimed to have produced strongly antitoxic plague sera by treating horses with filtrates from 8 to 10 weeks old bouillon cultures. He claims that 1 c. c. of his serum will neutralize 150 or 450 minimal lethal doses of the plague poison according to whether one measures the M L D by death in 48 hours or in 4 days. Rowland⁸¹ also has produced a serum by the immunization of animals with the "toxins" produced by his sulphate process. Rowland has apparently utilized the idea previously advanced by Lustig of immunizing with "nucleoproteins" derived from the plague bacillus instead of with the whole bacteria. Lustig's⁸² method consisted of washing up agar cultures of plague bacilli in 1 per cent. sodium hydrate solution, precipitating with ascitic acid, taking up the precipitate in an indifferent fluid and injecting it into horses. The serum produced by Lustig's method was used in Bombay, and is reported by Hahn as effective in milder cases, but without action in the more severe ones. There was but slight difference in the latter type between the treated and the untreated cases.

Rowland's⁸³ method consisted in the treatment of the moist bacteria with enough anhydrous sulphate of soda to combine with all the water present, freezing and thawing the mixture and filtering off the bacterial deposit at 37° C. Subsequently he extracted this bacterial mass with water. The extract so obtained was fatal to rats in quantities of 0.05 to 0.1 mg., killing them in 18 hours. In his experiments doses of 0.001 to 0.01 afforded protection, the last-named quantity reducing the mortality after inoculation of fatal doses from 80 per cent. to 10 per cent.

The sera produced by the immunization of horses with these supposed nucleoproteids are taken to be antitoxic in nature by Rowland himself and by MacKonky. They were used in the treatment of plague cases in the epidemics of 1908 and 1911 by the Maratha Hospital in Bombay, and reported upon by the Indian Plague Commission on the basis of observations made by Dr. Choksy. The cases in this series were controlled, as were those treated by the Yersin serum, by blood culture. Here the results were not striking—68.40 per cent. of the serum-treated cases died, while 77.60 per cent. of the controls died.

Altogether we cannot draw any definite conclusions as to the value of the serum treatment in plague. On the whole it does appear

⁸⁰ Dean. Cited from MacKonky, *Journ. of Hyg.*, Vol. 12, Plague Suppl. II, 1912, p. 402.

⁸¹ Rowland. *Journ. of Hyg.*, Vol. 11, Plague Suppl. I, pp. 11-19.

⁸² Lustig. "Monograph Sierotrapia e Vaccin Prev. Control la Peste," Turin, 1899; cited from Kolle and Martini, *loc. cit.*

⁸³ Rowland. *Journ. of Hyg.*, Vol. 10, p. 536.

that the milder cases are materially benefited by the treatment, and it is not at all impossible that in such cases aggravation of a milder case into fatal septicemia may be prevented by the timely administration of the plague serum. Animal experimentation also seems to indicate that the administration of the serum may be of great value as a prophylactic measure. It seems, on the other hand, as far as we can judge from the evidence of statistics, that when a case of plague has developed into the condition of active septicemia the administration of even the strongest plague sera at present available is of little use. And this is indeed unfortunately true of all passive immunization where the activity of the serum seems to depend chiefly upon bactericidal and opsonic properties. For we cannot definitely accept at the present day the claims that a true antitoxic serum, in the sense of those produced against diphtheria and tetanus poisons, can be really produced in the case of plague. The toxic substances derived from plague bacilli by a number of observers do not correspond in many particulars to true toxins.

FACTS CONCERNING ACTIVE PROPHYLACTIC IMMUNIZATION IN MAN

In a previous chapter we have dealt with the treatment of infectious disease with emulsions of dead bacteria or vaccines. The discussion there was confined to the use of these substances in the case of developed disease in which the infectious agent had already gained a foothold in the body. Concerning this form of therapy much difference of opinion exists, and we have seen that careful judgment must be applied to the selection of cases to which treatment with vaccines is adapted.

Concerning the *prophylactic* immunization of human beings with bacteria there can be little difference of opinion; this procedure finds its justification in prolonged laboratory experience in the hands of many men since the days of Pasteur.

The principle of specifically increasing the resistance of an individual by treatment with an altered form of the disease, either with the attenuated bacteria, with dead bacteria, or with bacterial extracts, has been sufficiently discussed in Chapter IV. It is indeed surprising that this phenomenon of prophylactic protection, though discovered by Jenner in small-pox, and developed by Pasteur in rabies, did not find more general application to the diseases of man until recent years. At present such methods are in extensive use in typhoid fever, in which they have had unquestionably excellent results. In the cases of cholera and plague numerous attempts have been made, but the results here are not as clear-cut as they have been in the case of typhoid. In the succeeding paragraphs we have set

forth as briefly as possible the methods employed in prophylactically immunizing man against this disease in which this procedure has been most commonly attempted.

PROPHYLACTIC IMMUNIZATION IN TYPHOID FEVER

The first attempt to inoculate human beings with typhoid bacilli with the intention of producing prophylactic active immunization was probably that made by Pfeiffer and Kolle ⁸⁴ in 1896. During the same year also Wright ⁸⁵ made similar studies in England, and soon after this he reported upon the development of antibodies in the blood of 17 people inoculated with typhoid. By these studies it was shown that human beings could be inoculated with dead typhoid bacilli without danger, and this logically led to the attempt to vaccinate human beings on a large scale.

It is hardly necessary to dwell upon the desirability of such a procedure. From tables recently published by Russell ⁸⁶ we take the information that, in our own Spanish-American war, 20,738 cases of typhoid with 1,580 deaths occurred in a total enlistment of 107,973. In this entire war 243 men were killed in action or died of their wounds, while almost 7 times as many died of typhoid fever. In the British army during the Boer war there were over 75,000 cases of typhoid in 380,000 men, and in the Russian army during the Russo-Japanese war over 17,000 cases of typhoid occurred, over half as many as the number of men killed in action. Such appalling figures leave no possible doubt as to the desirability of prophylactic immunization in armies, and there can be little question that typhoid fever is sufficiently prevalent in many parts of the civilized world to encourage prophylactic immunization of individuals, even when not living under the especially dangerous conditions of camps.

Following the preliminary studies of Pfeiffer and Kolle and of Wright extensive practical studies of vaccination were made in the German colonial army during the Herrero war, and by British bacteriologists during the Boer war. Leishmann ⁸⁷ also studied carefully the results of vaccination among regiments of the British army in India.

The vaccine employed by Wright and his associates in England consisted of broth cultures of a typhoid bacillus killed by exposure to 53° C., and by the further addition of 0.4 per cent. of lysol. The German vaccine consisted of emulsified agar cultures similarly killed.

The results obtained with these vaccines, although encouraging,

⁸⁴ Pfeiffer and Kolle. *Deutsche med. Woch.*, 22, 1896, p. 735.

⁸⁵ Wright. *Brit. Med. J.*, Jan., 1897, p. 256.

⁸⁶ Russell. *Amer. J. of Med. Sciences*, Dec., 1913, Vol. 146.

⁸⁷ Leishmann. *Glasgow Med. Journ.*, 1912, Vol. 77, p. 408, cited from Russell.

were not as striking as had been hoped. Russell⁸⁸ summarizes the earlier attempts by stating that the morbidity was reduced to about one half among vaccinated persons with a slightly greater reduction of mortality. The last-named writer also attributes the early failures to the overheating of the vaccines with a consequent reduction of their antigenic properties, and to timidity in their administration resulting from Wright's fear of a negative phase. Russell, of the United States Army Medical Corps, made a most extensive study of typhoid vaccination in this country. After careful consideration of the methods of others he produces his vaccines as follows: A single strain of typhoid bacilli is used (culture Rawlings obtained from England), and this is grown on agar in Kolle flasks for 18 hours. The purity of the culture is tested out both morphologically and by transplantation upon the double sugar media devised by Russell. Agglutination tests are also made. After 18 hours the growth is washed off in small quantities of salt solution and the emulsion heated in a water bath for one hour at 53° C.; it is then diluted with sterile salt solution to a concentration of one billion bacteria to a cubic centimeter. Then 0.25 per cent. of tricoresol is added. Before use the safety of the vaccine is ascertained both by aerobic and anaerobic cultivation and by the injection into mice and guinea pigs of considerable quantities for the exclusion of possible tetanus contamination. The efficiency of the vaccine is then tested by injecting rabbits with three doses at 10-day intervals, and determining the resulting agglutinating power.

With these vaccines under the direction of the United States Army Medical Corps the troops mobilized in Texas, California, and along the Mexican border were treated. Compulsory vaccination was established in March, 1911, and the results as reported by Russell have fully justified the measure. The following table taken from Russell's paper will illustrate the results obtained:

Typhoid Fever. Officers and Enlisted Men, United States Army

| | Yr. | Jan. | Feb. | Mar. | Apr. | May | June | July | Aug. | Sept. | Oct. | Nov. | Dec. | Totals for 9 months |
|------------|------|------|------|------|------|-----|------|------|------|-------|------|------|------|---------------------------|
| Voluntary | 1908 | 5 | 6 | 4 | 2 | 3 | 11 | 14 | 31 | 25 | 26 | 12 | 8 | 101 |
| | 1909 | 4 | 10 | 6 | 4 | 11 | 15 | 26 | 14 | 16 | 45 | 20 | 6 | 106 |
| | 1910 | 8 | 11 | 1 | 4 | 2 | 6 | 12 | 27 | 21 | 16 | 20 | 11 | 92 |
| Compulsory | 1911 | 3 | 3 | 3 | 7 | 4 | 4 | 4 | 7 | 4 | 4 | 1 | 0 | 39 |
| | 1912 | 1 | 2 | 2 | 0 | 0 | 3 | 1 | 3 | 1 | 4 | 0 | 1 | 13 |
| | 1913 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |

Paratyphoid fever included in figures for 1908, but excluded in other years. Cases paratyphoid, 1909, 3; 1910, 3; 1911, 2; 1912, 3; 1913, 0.

⁸⁸ Russell. *Am. Journ. of Med. Sc.*, Vol. 146, Dec., 1913.

We have mentioned in another place that Metchnikoff and Besredka in their studies on typhoid vaccination in the chimpanzee have concluded that very little protective value resided in vaccination with dead typhoid vaccines, whereas animals vaccinated with small amounts of living cultures were very efficiently protected. Metchnikoff and Besredka adopted finally the method of immunizing with living sensitized vaccines. By this is meant typhoid bacilli that have been exposed to the action of heated immune serum, or, in other words, typhoid bacilli that have absorbed specific antibodies. There is no question as to the efficiency of this form of vaccination. The method of employing sensitized bacteria for these purposes utilized by Besredka in the case of plague has unquestionably won an important place in active immunization. However, the results of Russell and others seem to indicate that in human beings the use of dead vaccines is certainly of considerable value, and there are certain practical objections to the use of living vaccines in immunization of large numbers of people as in armies to which Russell calls attention. In the first place, living vaccines cannot be stored for any considerable period, and may become a source of possible infection by mouth if carelessly handled; furthermore, contamination is not so easily ruled out in the case of living vaccines when used on a large scale, and it is not possible at present to require compulsory vaccination with living bacteria.

Gay has recently recommended the use of sensitized killed vaccines. He controls the efficiency of his vaccines by testing them out on rabbits in which typhoid septicemia has been produced by inoculation with cultures grown on rabbit blood agar. These vaccines have not yet been used upon sufficient numbers to justify conclusions. It would seem, however, that any one of the methods mentioned must possess considerable value, since they all represent merely slight variations of the same procedure. The method at present used in the German, British, and American armies, namely, vaccination with dead cultures, seems certainly, according to Russell's statistical studies, to have yielded excellent results and recommends itself by its extreme simplicity and safety.

ACTIVE PROPHYLACTIC IMMUNIZATION IN CHOLERA

Attempts to protect human beings against cholera by prophylactic vaccination were made as early as 1885 by Ferran,⁸⁹ a pupil of Pasteur. At the time at which Ferran's experiments were done little was known regarding the production of immunity with killed cultures or with bacterial extracts, and Ferran, under the influence of the French school and its endeavors to immunize with living attenu-

⁸⁹ Ferran. *C. R. de l'Acad. des Sc.*, 1885.

ated organisms, applied similar methods to cholera. First experimenting with guinea pigs, he soon applied his method to human beings, inoculating them with small quantities of living broth cultures of cholera spirilla. In many of his experiments he gave, at the first injection, 8 drops of a fresh broth culture, following this after 8 days with 0.5 c. c. of a similar culture. There is no reason why Ferran's method should not have yielded excellent results. However, it is stated that he worked with impure cultures, and other observers, notably Nikati and Rietsch, van Ermengen, da Lara, and others, failed to obtain encouragement in their subsequent investigation of this method of vaccination.

The method which Haffkine⁹⁰ worked out some years after Ferran's experiments also depended upon the injection of living cultures, but Haffkine attempted, by a rather elaborate technique, to produce two separate vaccines, one attenuated, the other enhanced in virulence. Attenuation was accomplished by growing the cholera spirilla at a temperature of 39° C. in broth over the surface of which a constant stream of sterile air was passed. Under these conditions the first crop of cholera organisms died rapidly, but Haffkine practiced reinoculation into new broth flasks before complete death of the original culture had taken place; after a series of generations of cultivation in this way he obtained cultures which produced merely temporary and slight edema when injected under the skin of guinea pigs. This weakened virus was used for the first inoculation.

He enhanced the virulence of cholera cultures with the purpose of producing a strain of maximum potency comparable to virus fixe. His procedure was as follows:

- a. Giving an animal an intraperitoneal injection of cholera spirilla larger than the fatal dose.
- b. Taking out the peritoneal exudate and exposing it for a few hours to the air.
- c. Injecting this exudate into another animal and treating the resulting peritoneal exudate in the same way.

After a series of such animal passages he claims to have obtained a virus of great virulence, and this is his second and stronger vaccine.

In applying the method to human beings Haffkine planted the cholera spirilla upon agar slants of the standard size, emulsified the growths in sterile water, and injected 1/5 to 1/20 c. c. of such a culture, using first the weak vaccine and five days later a more virulent culture.

Beginning his work as early as 1893, Haffkine and others vaccinated as many as 40,000 people in India. On the whole, the results obtained were very encouraging. It is a question, however, whether or not his method is unnecessarily complicated. In the light of our

⁹⁰ Haffkine. *The Lancet*, February, 1893; *Brit. Med. Journ.*, December, 1895.

more recent knowledge concerning cholera immunity it seems likely that the importance which Haffkine attached to the virulence of the cholera culture used for injection was exaggerated, and we have reason to believe that simple immunization with killed cultures may produce results fully as efficacious. After all, we could not expect, at least at present, to produce by active artificial immunization an immunity as permanent as that which results from an attack of the disease. Concerning the reasons for the acquisition of such permanent immunity we have as yet little knowledge. Even Haffkine's method of inoculation with living virus does not, by his own estimation, last longer than possibly two years. It is therefore likely that prophylactic immunization in cholera is efficacious by reason of the appearance in the blood serum of the specific bactericidal and opsonic substances by which the small numbers of cholera organisms entering during spontaneous infection can be disposed of before a foothold in the body is gained.

Tamancheff later used Haffkine's method, but killed the cultures by the addition of a 0.5 per cent. solution of carbolic acid.

Kolle⁹¹ later recommended the injection of dead cholera organisms, maintaining that a single injection of about 2 milligrams of a culture killed by exposure to 50° C. for a few minutes, and by the addition of 0.5 per cent. of phenol, is sufficient to immunize successfully. Good results with Kolle's method have been reported from Japan.

Strong,⁹² also proceeding from the idea that the immunizing antigen is present, as such, within the cell body of the cholera spirilla, recommends the injection of autolytic products obtained by digesting cholera spirilla in aqueous suspension and filtering. He prepared his "prophylactic" by growing the organisms upon agar, then suspending the growth in sterile water and keeping it at 60° C. for from one to twenty-four hours. The mixture was then exposed to 37° C. for from two to five days and filtered through Reichel filters. One to 5 c. c. of this was used in his experiments upon human beings.

PROPHYLACTIC IMMUNIZATION AGAINST PLAGUE

The first attempts to immunize human beings prophylactically against plague were those of Haffkine.⁹³ The first vaccinations were carried out with broth cultures killed at 65° C. He tested out his vaccines on a large scale in Bombay, and obtained apparently promising results. In a plague epidemic occurring in a Bombay prison

⁹¹ Kolle. *Deutsche med. Woch.*, 1897, No. 1.

⁹² Strong. *Journ. Inf. Dis.*, Vol. 2, 1905.

⁹³ Haffkine. *Bull. de l'Inst. Past.*, Vol. 4, 1906, No. 20, p. 825.

only 2 of 151 vaccinated persons became ill, and neither of these died; whereas, of 177 unvaccinated persons 12 became ill and 6 died. In large series of vaccinated people only 1.8 per cent. were infected with plague, with a mortality of 0.4 per cent. for the total, whereas of unvaccinated individuals in the same epidemic 7.7 per cent. fell victim to the disease, with a mortality of 4.7 per cent.

The German Plague Commission, consisting of Gaffky, Pfeiffer, and Dieudonné, recommended a vaccine of killed agar cultures. Kolle and Otto,⁹⁴ basing their earlier results upon experiments carried out with monkeys, mice, guinea pigs, and rats, have come to the conclusion that vaccination with dead plague cultures is much inferior to that obtained when attenuated living cultures are used. The same conclusion has been reached by Kolle and Strong.⁹⁵ Kolle and Otto found that the immunization of animals with large doses of killed agar cultures of plague bacilli and with Haffkine's prophylactic did not protect them against subsequent inoculation with virulent cultures.

Strong⁹⁶ subsequently made a very careful comparative study of the various methods of plague vaccination, and concluded that the most efficient method is immunization with attenuated living cultures. He showed that when carefully done this method can be safely employed in human beings, but admits that his work must be as yet considered as experimental and further studied before it can be universally employed.

Besredka⁹⁷ has advised the use of sensitized dead plague cultures, claiming, from animal experimentation, that such vaccines produce an efficient and relatively durable immunity.

Rowland⁹⁸ confirms the immunizing properties of Besredka's vaccines in plague, and believes that the antigenic properties of the plague bacillus are attached to the bacterial nucleoproteins, and can be extracted with these. Rowland prepares a vaccine by the treatment of the moist bacteria with enough anhydrous sodium sulphate to combine with all the water present, freezing and thawing the mixtures, then filtering off the bacterial deposits at 37° C., and extracting them with water. The solution so obtained was fatal to rats in small quantities and afforded substantial protection, reducing the mortality on subsequent inoculation of a standard culture from 80 to 10 per cent.

⁹⁴ Kolle and Otto. *Deutsche med. Woch.*, 1903, p. 493, and *Zeitschr. f. Hyg.*, Vol. 45, 1903.

⁹⁵ Kolle and Strong. *Deutsche med. Woch.*, XXXII, 1906, p. 413.

⁹⁶ Strong. *Journ. of Med. Res.*, N. S., 13, 1908.

⁹⁷ Besredka. *Bull. de l'Inst. Past.*, Vol. 8, 1910.

⁹⁸ Rowland. *Journ. of Hyg.*, Vol. 12, 1912, p. 344.

PROPHYLAXIS AGAINST SMALL-POX

In the case of small-pox the general method of active prophylactic immunization is in principle identical with that devised by Jenner in the 18th century. The original observation from which Jenner worked was that dairy maids and other individuals who had been infected with cow pox were thereafter spared when a small-pox epidemic appeared in the region in which they lived. It is now agreed by most observers who have studied the problem that the virus of cow pox and that of small-pox are identical in nature; the former representing a strain attenuated by passage through the animal body. This is based chiefly upon the observation that true variola can be transmitted to cattle, and that it can be thus carried from animal to animal, during this process becoming attenuated for human beings to such a degree that reinoculated into man a simple vaccinia is produced.⁹⁹

Small-pox, therefore, represents in principle active immunization by means of attenuated virus. When vaccination was first introduced the virus was taken from preceding pustules produced in other human beings. This has been given up in most countries to-day largely because of the dangers of transferring syphilis and other diseases in this way. At present the method of obtaining virus for vaccination purposes is carried out as follows: The initial material consists of what is known as "seed" virus, which can be obtained from spontaneous cow pox or from vaccination pustules in children, or again from pustules obtained in calves after several passages of small-pox virus through these calves. From such seed virus calves may be inoculated for vaccine production or else the calves may be inoculated from the material obtained from other calves in the usual way.

Healthy young animals are used; they are washed along the abdomen, strapped down upon specially prepared tables, and the abdominal skin thoroughly cleansed with soap and water. The exact procedure varies in different places; often the skin is thoroughly cleansed with carbolic solution, and this is thoroughly removed with sterile water before inoculation, or else cleansing is relied upon without the use of germicides. Over the clean area longitudinal scratches 1 to 2 c. c. apart are made, and into these the seed virus is rubbed. The animals are then kept in a clean stall, preferably over asphalt floors, and rigid cleanliness is observed during the period of development of the pustules. After the 6th or 7th day, when the vesicles are beginning to appear, the abdomen is well washed and cleansed of superficial dirt without the use of an antiseptic, and the pulp removed from the lesions with a curette. The pulp so removed is placed into 60 per cent. glycerin and thoroughly ground up in a specially constructed mill. According to Rosenau, the animal should always be killed before the vesicles are

⁹⁹ Haccius. Cited from Paul, Kraus and Levaditi, Vol. 1, p. 593.

removed, not only for humane reasons, since the same object might be attained with anesthesia, but because a thorough autopsy can then be performed to determine the health of the calf.

Vaccines so obtained always contain bacteria, the glycerin therefore serving a double purpose: one, the preservation of the virus, the other a gradual destruction of the bacteria. Rosenau has shown that the addition of 2 to 4 parts of 60 per cent. glycerin to one part weight of the pulp prevents the growth of bacteria and probably destroys them by dehydration. Most of the bacteria are destroyed within one month at 20° C. During this period, then, from 4 to 6 weeks, the glycerinated virus should not be used, and should from time to time be controlled by cultivation. At the end of this time the lymph is ready for use.

Formerly the material for the vaccination of human beings was obtained very simply by dipping ivory splinters into the fluid of pustules, allowing this to dry, and rubbing these ivory or bone points into the exudate obtained by scratching the skin of the individual to be vaccinated. This method has practically gone out of use, and to-day the ripened glycerin pulp prepared as above is taken up in small capillary glass tubes and from these blown upon the vaccination scratch. The efficiency of vaccine virus can be tested for potency by the inoculation of the ears of rabbits before use.

ACTIVE PROPHYLACTIC IMMUNIZATION IN RABIES (HYDROPHOBIA)

Although many modifications have been suggested and actually used in different parts of the world, the most common method of immunizing against rabies still remains that originally devised by Pasteur. The Pasteur treatment takes advantage of the prolonged incubation period of rabies and is planned to confer immunity between the time of inoculation and the time at which the disease would naturally appear. Since this period in ordinary street infection by dog bite is usually 40 days or more, a considerable interval for active immunization is available. Formerly much of this time was lost in that the diagnosis of hydrophobia in the dog or other animal that had caused the injury could not be made with certainty until the results of rabbit inoculations had been obtained. Nowadays the ease with which a diagnosis of hydrophobia can be made within a few minutes by finding negri bodies in the hippocampal and cerebellar cells has added considerably to safety in that it has made possible a gain of almost two weeks in determining whether treatment should be instituted or not.

Here again, although the infectious agent of rabies is not known with certainty even at the present day, the method of Pasteur depends upon active immunization by means of an attenuated virus.

In standardizing the virus for the purpose of treatment Pasteur first produced what he calls the "virus fixe." This consists of the

ordinary street virus as obtained from rabid animals passed through a considerable series of rabbits (20-30) until its virulence for these animals has reached a maximum. After a sufficient number of such rabbit passages the incubation time after intracerebral inoculation is reduced to 7 or 8 days, but can no longer be shortened by further passage. The brain and cord material of rabbits dead of rabies after such repeated passages constitutes virus fixe. This can be preserved for considerable periods in 60 per cent. glycerin, and this is the initial material from which the attenuated preparations for treatment are produced.

In preparing the material for treatment a small amount of virus fixe is injected subdurally into rabbits, about 0.2 c. c. of a salt solution emulsion being given. The inoculation is very easily made through a small trephine opening in the skull, and contamination is very easily avoided. Just before the rabbit dies when completely paralyzed it is killed by chloroform and the cord is removed best by the method of Oschida.¹⁰⁰ The rabbit is nailed to a board, back uppermost, and washed with a weak antiseptic, a longitudinal incision is then made along the backbone from the occiput to the lumbar region, and the vertebral column laid bare.

After searing the tissues around the back of the head the spine is cut across just behind the occiput, and again in the same way just above the sacrum. The neck and lumbar regions are dissected loose from the skin and gauze is inserted under it to avoid contamination. The assistant grasps the end of the spinal cord as it appears in the cervical region and pulls on it very slightly while the operator with a glass rod or a piece of wire pushes against it from below. If this is carefully done the spinal nerves are torn and the cord can be gradually pulled out of place. This procedure is by far the best, although it requires a certain amount of practice.

The cords so removed are hung up by a thread in bottles containing sticks of caustic potash and exposed in a dark place to 22° to 23° C. Under these conditions of drying and temperature the virus is gradually attenuated until at the end of 13 days or more the virulence is practically nil. If removed from the drying bottles at any time during the process and kept in a refrigerator in sterile glycerin the virulence, whatever it may be at the time of placing into the glycerin, remains fairly constant for long periods. When any of this material is used for treatment little pieces of the cord $\frac{1}{8}$ cm. in length are cut off and emulsified in 2.5 c. c. of salt solution, and this emulsion is used for injection.¹⁰¹

¹⁰⁰ Oschida. *Centralbl. f. Bakt.*, Vol. 29, 1901.

¹⁰¹ In our description of the methods of drying rabies, for the sake of adhering to a standard, we follow closely the directions laid down by A. M. Stimson, in the *U. S. P. H. S. Bull.* 65, 1910. There are various modifications used in different countries, in many cases unimportant, and it seems well to adhere to the U. S. regulations as a standard for this country.

When patients are to be treated the principle of the treatment is to inoculate them first with cords that have been dried for considerable periods, gradually proceeding toward those that have been dried for less prolonged times and are therefore more virulent. The treatment is varied in the individual case according to the severity of the injury. Formerly treatment was begun with cords dried as long as 16 days. More recently it has been found that cords dried for longer than 8 days are practically non-virulent and correspondingly lack in antigenic value. They are no longer employed therefore, since their use is regarded as a waste of time. The following tables taken from Stimson's article in Bulletin 65 of the Hygienic Laboratory of the U. S. Public Health Service give the standard methods of treatment as recommended by the United States Public Health Service:

Scheme for Mild Treatment

| Day | Cord (injections) | Amount injected | | | Day | Cord (injections) | Amount injected | | |
|-----|----------------------|------------------|-------------------------|------------------------|-----|----------------------|------------------|-------------------------|------------------------|
| | | Adult (c. c.) | 5-10 yrs. (c. c.) | 1-5 yrs. (c. c.) | | | Adult (c. c.) | 5-10 yrs. (c. c.) | 1-5 yrs. (c. c.) |
| 1 | 8-7-6=3 | 2.5 | 2.5 | 2.0 | 12 | 4=1 | 2.5 | 2.5 | 2.5 |
| 2 | 5-4=2 | 2.5 | 2.5 | 1.5 | 13 | 4=1 | 2.5 | 2.5 | 2.5 |
| 3 | 4-3=2 | 2.5 | 2.5 | 2.0 | 14 | 3=1 | 2.5 | 2.5 | 2.0 |
| 4 | 5=1 | 2.5 | 2.5 | 2.5 | 15 | 3=1 | 2.5 | 2.5 | 2.0 |
| 5 | 4=1 | 2.5 | 2.5 | 2.5 | 16 | 2=1 | 2.5 | 2.0 | 1.5 |
| 6 | 3=1 | 2.5 | 2.5 | 2.0 | 17 | 2=1 | 2.5 | 2.0 | 1.5 |
| 7 | 3=1 | 2.5 | 2.5 | 2.0 | 18 | 4=1 | 2.5 | 2.5 | 2.5 |
| 8 | 2=1 | 2.5 | 1.5 | 1.0 | 19 | 3=1 | 2.5 | 2.5 | 2.5 |
| 9 | 2=1 | 2.5 | 2.0 | 1.5 | 20 | 2=1 | 2.5 | 2.5 | 2.0 |
| 10 | 5=1 | 2.5 | 2.5 | 2.5 | 21 | 2=1 | 2.5 | 2.5 | 2.0 |
| 11 | 5=1 | 2.5 | 2.5 | 2.5 | | | | | |

Scheme for Intensive Treatment

| Day | Cord (injections) | Amount injected | | | Day | Cord (injections) | Amount injected | | |
|-----|----------------------|------------------|-------------------------|------------------------|-----|----------------------|------------------|-------------------------|------------------------|
| | | Adult (c. c.) | 5-10 yrs. (c. c.) | 1-5 yrs. (c. c.) | | | Adult (c. c.) | 5-10 yrs. (c. c.) | 1-5 yrs. (c. c.) |
| 1 | 8-7-6=3 | 2.5 | 2.5 | 2.5 | 12 | 3=1 | 2.5 | 2.5 | 2.0 |
| 2 | 4-3=2 | 2.5 | 2.5 | 2.0 | 13 | 3=1 | 2.5 | 2.5 | 2.0 |
| 3 | 5-4=2 | 2.5 | 2.5 | 2.5 | 14 | 2=1 | 2.5 | 2.5 | 2.0 |
| 4 | 3=1 | 2.5 | 2.5 | 2.0 | 15 | 2=1 | 2.5 | 2.5 | 2.0 |
| 5 | 3=1 | 2.5 | 2.5 | 2.0 | 16 | 4=1 | 2.5 | 2.5 | 2.5 |
| 6 | 2=1 | 2.5 | 2.0 | 1.5 | 17 | 3=1 | 2.5 | 2.5 | 2.5 |
| 7 | 2=1 | 2.5 | 2.5 | 2.0 | 18 | 2=1 | 2.5 | 2.5 | 2.0 |
| 8 | 1=1 | 2.5 | 1.5 | 1.0 | 19 | 3=1 | 2.5 | 2.5 | 2.0 |
| 9 | 5=1 | 2.5 | 2.5 | 2.5 | 20 | 2=1 | 2.5 | 2.5 | 2.5 |
| 10 | 4=1 | 2.5 | 2.5 | 2.5 | 21 | 1=1 | 2.5 | 2.5 | 2.0 |
| 11 | 4=1 | 2.5 | 2.5 | 2.5 | | | | | |

This is the standard treatment used almost everywhere in the world at present. Other methods have been recommended. One of these is that of Högyes, in which virus fixe unattenuated is used in dilution. Högyes begins by injecting 3 c. c. of a 1 to 10,000 dilution of virus fixe, gradually proceeding within 14 days to 1 c. c. of a 1 to 100 dilution.

Fixed virus attenuated by the addition of antirabic serum and chemical disinfectants (carbolic acid) and by partial digestion in gastric juice has also been used, but none of these methods has attained widespread application.

CHAPTER XX

ABDERHALDEN'S WORK UPON PROTECTIVE FERMENTS OF THE ANIMAL BODY

THE recent researches of Abderhalden¹ upon the intravascular digestion of foreign substances introduced into animal bodies promise to have considerable bearing upon problems of immunity. Abderhalden, whose work we cite chiefly from his monograph, "Die Schützfermente des tierischen Organismus," took as his point of departure the conception that the animal body must necessarily dispose over a mechanism whereby it can assimilate foreign substances which obtain entrance unchanged into the circulation. In our section upon the nature of the precipitins, especially in the discussion of Gengou's conception of "albuminolysins," we have called attention to the probable significance of protein antibodies as a mechanism for the disposal of such foreign substances. In the bodies of the higher animals in which a special alimentary system, with its many digestive ferments, is well developed, it is most probable that the normal condition of digestion is one in which the foreign substances utilized for nutrition are completely split into their simpler components before they gain entrance to the circulation. Nevertheless, abnormal conditions or accidents, such as gastro-enteric diseases, digestive disturbances, and bacterial infections, may lead to a condition, probably frequent enough in ordinary life, during which such foreign substances may get into the blood stream without previous cleavage. The problem is to determine where and how such substances, protein or otherwise, are broken up so that they may be either assimilated or eliminated. We have referred in another place to the fact that foreign proteins may occasionally pass through the kidneys and be eliminated unchanged. This has been shown actually to occur by Oppenheimer, Ascoli, and others, but probably represents a very unusual state of affairs produced by special experimental conditions. As a rule these substances are disposed of within the body by chemical cleavage or by assimilation. Abderhalden believes that this process depends upon the mobilization of "protective ferments," a term which he borrows from Heilner,² and suggests the possibility

¹ Abderhalden. "Schützfermente des tierischen Organismus," Springer, Berlin, 1912.

² Heilner. Cited from *Abderhalden Zeitschr. f. Biol.*, Vol. 50, 1907.

that these ferments may possibly originate in the leukocytes. He refers to the work of Friedrich Müller, in which it was shown that the resorption of pneumonic consolidations is largely carried on by leukocytic ferments. Moreover, we possess in support of such a conception the many consistent reports of the successful extraction of various ferments from leukocytes, some of which are referred to in detail in another section.

Experimentally Abderhalden approaches his problem by determining the presence of specific ferments in the blood of animals into which various foreign substances have been introduced by paths other than the alimentary canal. For this purpose he has developed a number of methods, the most important of which are his optical method and his dialysis method. The optical method used for the determination of the proteolytic properties of the serum depends upon the fact that many of the amino-acids are optically active. Moreover, most of these substances are chemically known and their optical activity determined, so that it is possible to take blood serum which is to be examined for its contents of particular ferments, mix them with a suitable protein, or preferably a polypeptid, and determine with a polariscope the rotation which takes place. We will not go into the technique of this method more extensively because we have no personal experience with it, and the method is one of such delicacy that it is best obtained from Abderhalden's original publications directly.³ His dialysis methods depend upon placing the blood serum and fermentable substance into dialyzing bags, suspending them into distilled water, and determining the presence of peptone, amino-acids, or total nitrogen in the liquid outside of the bag after definite intervals of time.

By these and other methods Abderhalden⁴ has carried out tests with a large number of different substances. Experimenting first with proteins, he injected egg albumen, horse serum, silk peptone, gelatin, edestin, casein, etc., into dogs and rabbits, then, several days later, bled the animals and mixed 0.5 c. c. of the serum with 0.5 c. c. of a solution of the respective substances which had been injected. He found in such cases that definite proteolytic action was exerted upon the injected substances by the active serum of a treated animal, whereas, in the case of most of the substances used, the normal serum possessed no proteolytic action whatever. These results were consistently obtained both by the dialysis and by the optical methods. It should be especially noted that the ferments studied by Abderhalden were not as specific as are the antibodies which we have discussed in another place. For Abderhalden found that the serum of

³ See especially Abderhalden, Hoppe-Seyler, *Zeitschr. f. physiol. Chemie*, Vols. 60, 65, and 66; also "Handbuch der biochem. Arbeitsmethoden," Vol. 5, p. 575, 1911.

⁴ Abderhalden. "Schützfermente," p. 49.

an animal treated with proteins developed enzymes which were active, not only against the particular protein used for injection, but rather against proteins in general. They were specific only in that, when produced with proteins, they were not active against fats or carbohydrates. This is especially important in connection with the recent discussion concerning the identity of Abderhalden's protective ferments and the specific protein antibodies.

In later experiments Abderhalden showed further that similar ferments could be induced in animals by treatment with carbohydrates and with fats. The serum of normal dogs is not capable of splitting cane sugar. However, the blood serum or plasma of a dog that has been treated with cane sugar develops the property of inverting the cane sugar into dextrose and fructose within fifteen minutes after injection. This could easily be determined both by putting together the serum with cane sugar and determining the increase of reducing powers, and by means of subjecting such active plasma or serum, together with saccharose, to polariscopic examination.

The earlier experiments with fats were negative because the simple method of titration for fatty acids proved insufficient as an indicator of activity. However, Abderhalden succeeded in determining fat-splitting properties in the blood of treated dogs by using the method of Michaelis and Rona.⁵ The presence of fats largely increases the surface tension of mixtures, and their cleavage in such mixtures consequently leads to reduction of this tension. Utilizing this principle, Abderhalden claims to have determined that the parenteral introduction of fats into dogs is followed by a reactionary increase of lipases.

The general significance of Abderhalden's researches is this: When any foreign substances, protein, carbohydrate, or fats, gain entrance to the circulation of an animal, the animal body reacts by the mobilization of ferments or enzymes specifically capable of reducing these substances to assimilable form. It is likely that these ferments represent a mobilization of substances normally present but not concentrated in the blood stream under ordinary conditions, since they appear with a speed out of all proportion to that obtaining in the case of the antibodies discussed in another place. In one case cited by him a dog injected on November 25th, 29th, and December 4th showed powerful peptolytic serum properties on December 6th. Apparently the injection of homologous proteins into animals (i. e., rabbit serum into rabbits, etc.) does not incite reaction.

These enzymes seemed to differ from specific antibodies in that they did not react solely with the substance injected, but also with other substances belonging to the same chemical group. Other differences from antibodies are the rapid appearance of the ferments

⁵ Michaelis and Rona. Cited from Abderhalden, *loc. cit.*

after treatment and their rapid disappearance after the inciting stimulus is removed. Thus Abderhalden reports that the enzymes found in a case of pregnancy disappeared within eight days after abortion or child birth.

It is plain that these researches of Abderhalden offer many opportunities for diagnostic utilization, and he has applied them to the diagnosis of pregnancy. In this condition substances from the chorionic villi get into the blood. These, according to Abderhalden, may be looked upon as in a certain sense foreign in nature, and must be chemically disintegrated by the body. In consequence it is likely that the ferments which accomplish this would appear in the sera of pregnant individuals and could be determined by his methods. When he prepared peptone from the placental substances of human beings and allowed the blood plasma of normal individuals to act upon it, observing it both by the dialysis and the optical method, no peptolytic action could be observed. However, when the plasma of pregnant women was used proteolytic action was determined. In these cases the ferment seemed to be specific for peptones produced from placental tissue both in animals and human beings, but did not act upon casein, gelatin, or other proteins. There are certain technical difficulties connected with the production of a test material from the placental tissue which render this method difficult. For their more detailed description we refer the reader to the original articles. Abderhalden believes that his protective ferments may have considerable bearing upon the problems of bacterial immunity and anaphylaxis, and this of course is evident to every one who has followed the development of these subjects. The problem, however, is a complicated one, and it is quite impossible at present to draw definite conclusions.

THE MEIOSTAGMIN REACTION

Ascoli and Izar⁶ have attempted to work out a diagnostic reaction which depends upon an alteration of surface tension of a fluid when an antigen unites with its specific antibody. Ascoli in his first experiments worked with typhoid bacillus extracts and the sera of typhoid patients, and found that when the two suspensions were mixed a reduction of surface tension resulted after time for union between the two had been allowed.

They determined the reduction of surface tension by Traube's⁷ method by the use of apparatus spoken of as the "stalagmometer." The principle of this method depends upon the fact that as surface tension is reduced the number of drops to a given quantity of fluid is increased.

⁶ Ascoli and Izar. *Münch. med. Woch.*, Nos. 2, 7, 18, 22, 41, 1910.

⁷ Traube. *Pflüger's Archiv*, Vol. 123, 419.

Diluted serum of patients was mixed with diluted antigen, and the number of drops contained in one cubic centimeter of the mixture was immediately determined and again measured after the mixture had remained for two hours in the incubator at 37° C. An example of one of Ascoli's early measurements is given in the following protocol:

1 c. c. of serum of typhoid patient diluted to 1-10.

| | | Number of drops | |
|---|----------|------------------|----------------------------------|
| | | Immedi- ately | After 2 hours in incubator |
| 1 c. c. alcoholic typhoid extract diluted to... | 1 0/00 | 57.8 58.1 | 59.7 59.9 |
| | 1 0/000 | 57.5 57.5 | 59.4 59.6 |
| | 1 0/000 | 57.0 57.0 | 59.3 59.2 |
| | 1 0/00 | 58.1 57.7 | 59.7 59.6 |
| | 1 0/000 | 57.4 57.6 | 59.4 59.2 |
| | 1 0/000 | 57.0 56.9 | 59.2 59.4 |
| 1 c. c. alcoholic precipitate taken up in distilled water. | 1 0/00 | 56.5 56.5 | 58.0 57.8 |
| | 1 0/000 | 56.5 56.6 | 57.5 57.4 |
| | 1 0/0000 | 56.7 56.5 | 57.4 57.5 |
| | | | |
| 1 c. c. in 1 0/00 alcohol in 1 c. c. 0.85 per cent. NaCl solution..... | | 56.6 56.7 | 57.5 57.6 |

Of course a certain amount of reduction of surface tension results when various antigens are brought together with normal sera, but this can be easily controlled by suitable dilution, and must be carefully taken into consideration in each individual case. Ascoli and Izar have applied this method to the diagnosis of tuberculosis, typhoid, and various other diseases, and have reported what seemed to them reliable results. So far experience with the meiostagmin reac-

tion has not been very extensive; not all observers have been able to obtain results as apparently reliable as those of Ascoli and his collaborators. It is not possible therefore to express a final opinion regarding this method of investigation; it contains, however, an interesting principle which with more exact methods of measurement may well become very important in serum diagnosis.

CHAPTER XXI

COLLOIDS

BY STEWART W. YOUNG

Professor of Physical Chemistry, Stanford University, Cal.

INTRODUCTORY

IN attempting to give in the brief space of a single chapter any adequate account of the present state of our knowledge in so vast a field as that of colloid chemistry and physics one is confronted with a rather difficult problem. In the present outline the attempt will be made to get at some notion of the matter by a presentation first of the more important generalizations which have been drawn, this to be followed in each case by sufficient experimental evidence to serve as illustration, together in some cases perhaps with certain evidence which may seem to contradict in some degree such current conceptions. The reason for this particular method of presentation lies in the fact that new material is so rapidly accumulating, much of which seems more or less at variance with present accepted theories that it seems more than possible that some of these fundamental generalizations may soon undergo material modification if not reform. It would, therefore, seem ill-advised, in presenting a brief résumé to readers who are not physicists or chemists, merely to present the present theories as they are used to-day by workers in the field, and to sound a note of warning that many, if not all of them, are not so securely supported by broad evidence as to allow of very concrete prediction being based upon them.

Definition.—The fundamental distinction between the crystalloid and colloid states of matter was first drawn by Thomas Graham¹ as a result of his investigations into the phenomenon of dialysis. He noted that in general those substances which when in solution did not pass through the dialyzing membrane, or did so only very slowly, also were characterized by the fact that when they separated from solution, either by precipitation or by evaporation, they did so in the non-

¹ Graham. *Phil. Trans.*, 1861, 183.

crystalline or amorphous form. This class of bodies he named colloids, since glue (Greek *κολλα* meaning glue) presented a typical case. Colloid substances may appear in highly dispersed states, such as dilute glue, arsenic sulphid suspensions, oil or rosin emulsions, milk (casein in highly dispersed condition), and the like, in which case they are spoken of as *sols*. They may also occur in the undispersed or only slightly dispersed state, as the amorphous precipitated sulphids of the heavy metals, precipitated casein, or dry glue. In this state they are spoken of as *gels*.

When a colloid substance has once been converted from the *sol* or dispersed state into the *gel* or undispersed state, its properties may differ greatly in different cases. Thus if a dispersed soap (soap solution, or more correctly soap sol) be coagulated by the addition of common salt, the coagulum or soap gel may be removed from the salt solution, and if again placed in pure water it will redisperse and again assume the sol condition. Such a colloid is spoken of as a *reversible colloid*. If, however, an arsenious sulphid suspension be put through precisely the same course of treatment it will, in the last stage of the treatment, refuse to redisperse, and is therefore spoken of as an *irreversible colloid*. Some authorities prefer to speak of these two classes of colloids as emulsion and suspension colloids, respectively,² since in general those colloids which are reversible tend to separate out in soft masses, and in general to gelatinize rather than to flocculate, while the irreversible colloids rather tend to truly flocculate and form very compact and frequently more or less granular coagula. Since, however, we seem more likely at the present time to suffer more from an excess of classification than from a lack of it, the attempt will be made to get along in this discussion with the earlier nomenclature. It may, indeed, be added that it is highly probable that the distinction between reversible and irreversible colloids is only one of degree. For example, many of the metallic sulphids which are typically irreversible may be made to some extent reversible by means of thorough washing and re-treatment with hydrogen sulphid which had been originally used in their preparation. It is probable that certain colloids are apparently irreversible only because we do not truly reverse the conditions.

Heretofore in this discussion the term "colloid" substance has been used as if to imply that certain chemical individuals were characteristically colloid, while others were not. It was much in this sense that Graham used the term. Investigations since his time have shown this to be a misconception, and it is now apparent that any and all substances may be either colloid or crystalloid, the form they assume depending upon treatment. Thus albumin may be crystallized and common salt may be obtained in the state of a colloid solu-

² V. Weimarn. *Ztschr. Chem. Ind. Koll.*, 1908, 3, 26.

tion or sol.³ Albumin, gelatin, and agar may be obtained crystalline by proper regulation of temperature and the use of proper solvents, as solutions of ammonium sulphate for albumin, and alcohol-water mixtures of varying strengths for the two latter substances. Sodium chlorid has been obtained in the colloidal condition by precipitating it in a solution of sodium sulphocyanate by hydrochloric acid, each of the reacting substances being dissolved in a mixture of amyl alcohol and ethyl acetate.

There is much evidence that leads to the belief that all colloid systems are unstable. Van Bemmelen characterized them as systems which never reached a state of rest, that is, were never in equilibrium. The conditions which determine the appearance of a body in the colloid or crystalline form lead to the suspicion that bodies always separate from solution in the amorphous or colloidal condition and that all crystallization is a secondary phenomenon. The conditions that are favorable for the transformation of a colloid into a crystalline form are a considerable solubility and a considerable rate of crystallization. Where either or both of these is at a minimum the conditions are favorable for relative permanence in the colloid condition. It is upon the basis of this principle that von Weimarn succeeded in obtaining relatively stable colloidal solutions of common salt and many other easily crystallizable salts. Furthermore Doelter⁴ has succeeded in converting many well-known amorphous precipitates into crystalline bodies by means of stirring, pressure, impact, and high temperature. Among the substances thus transformed are aluminium, chromium, and iron hydroxids, and the sulphids of arsenic, antimony, and zinc.

With this much by way of introduction, we may now proceed to a closer consideration of some of the better recognized properties of colloid sols and gels. For convenience we shall first take up the discussion of these systems from the more definitely physical point of view, and later take up those properties which seem more definitely chemical.

Physical Properties of Colloids.—1. FORM AND SIZE.—Current opinion seems to be leading rapidly to the general acceptance of the hypothesis that in liquid systems of two or more components we have to do with a continuous series of conditions ranging from coarse suspensions through suspensions of increasing fineness (increasing degrees of dispersion) to finally the molecular and ionic states of solution. The opinion is also growing that, although for certain practical purposes the classification of all such systems in one way or another, as in terms of the various degrees of dispersion, may be useful, the

³ V. Weimarn. *Ibid.*, 1910, 7, 92, and "Grundzüge der dispersoid Chemie," 107-108.

⁴ *Ztschr. Chem. Ind. Koll.*, 1910, 7, 86.

excessive use of such classifications is likely to narrow rather than broaden our conception of the whole subject matter of the field. It would seem that the most stimulating point of view is to be reached from the acceptance of the suggestion of Wolfgang Ostwald, that the chief problem of colloid chemistry at the present time lies in determining the influences of the degree of dispersion upon the physical and chemical properties of all liquid solutions, mixtures, suspensions, or what-not. If this point of view be taken it follows that the form and size of the particles in a disperse system are a matter of the first importance.

If the degree of dispersion in a given system be not too great the form of the particles may be readily observed under the microscope. Such evidence shows that the spherical form predominates enormously over all others, although under carefully controlled conditions ovoid forms may appear, as in the case of gelatin and agar. These ovoid forms are taken by von Weimarn (*loc. cit.*) and others, as evidence of directive forces, and hence of incipient crystallization. If the system be treated in such a way as to decrease the dispersion, as, for example, if a reagent be added which tends to flocculate the colloid, but not in sufficient quantity to produce actual precipitation, the decrease in dispersion may take place in two quite different ways: first, the size of the particles may increase, as in the case of oil emulsions; second, the particles do not coalesce but become attached together in chains and groups which, in many cases, resemble bunches of grapes. This sort of aggregation may go so far as to produce web-like structures. The jellying of gelatin has been shown to be due to the development of such web structures. Glue shows much less tendency in this direction, and if some acetic acid be added, as in the preparation of commercial liquid glues, this web formation is almost entirely absent, and the adhesive qualities are at the same time greatly improved. It seems quite certain that both of the above modes of aggregation are possible in one and the same system at different stages in its condensation. Thus highly dispersed copper sulphid becomes aggregated in its first stages of condensation by an actual increase in the size of the spherical particles. After these reach a certain fairly definite size further aggregation takes place by the grouping together of these spheres. It is generally recognized that all grouped and webbed structures are secondary.

A large number of very important investigations have been directed toward the determination of the size of disperse particles throughout the greatest variation in dimensions. The fact first noted by Graham, that substances in colloidal solution show a very small, and frequently almost negligible, rate of dialysis, points directly to the supposition that the particles in such solutions are in a far less dispersed state than in solutions of crystalloid substances. The rate of dialysis is directly determined by the rate of diffusion,

which, in turn, is inversely proportional to the square roots of the masses of the diffusing bodies.

Measurements of osmotic pressure in solutions also give an accurate measure of the relative masses of dispersed systems where such measurements can be successfully carried out, and a great deal of work has been devoted to attempts to measure the osmotic pressure of colloidal solutions. Great difficulties both of experimentation and of interpretation are encountered in this field. As will soon be seen a colloid particle stands in a very complex relationship to its surrounding liquid, and furthermore it is a matter of extreme difficulty to obtain a colloid solution free from electrolytes, which themselves may create osmotic pressure or otherwise affect the measurements. About the only conclusion which it is safe to draw at the present time is that if colloid solutions show osmotic pressure at all the value of it is very small compared to that shown by crystalloidal solutions of substances of more or less like formula weights. This leads to the conclusion that the particles in a colloid solution are in a state of dispersion far less than that found in a typical crystalloidal solution. For a most excellent résumé of the present state of our knowledge in this field the reader is referred to a recent book by Dr. L. Casuto, of Pisa, entitled "*Der Kolloide Zustand der Materie*" (Steinkopf, 1913).

When the size of the disperse particles is sufficiently great they may, of course, be measured under the microscope, and with the advent of the ultramicroscope the limits of visibility of small bodies has been very notably extended. The ultramicroscope is known in several forms, the first having been devised by Siedentopf and Zsigmondy. All depend upon the production of powerful rays of light in directions parallel to the surface of the microscope slide. In such a field there will be no luminosity, provided the field is optically empty, that is, contains no particles of sufficient size to produce a dispersion of light. If, on the other hand, such particles are present, the effect observed will be an illumination whose character will depend upon the size of the particles. If the particles are of sufficient size the illumination will show them individually as bright points of dispersion, even though the particles are too small to be observed of themselves, just as the stars are visible from the light which they disperse, but cannot of themselves be seen. If the particles are so small that they are no longer able to disperse sufficient light to make each particle appear as a bright point, there will, nevertheless, provided the particles are present in sufficient numbers, be produced a diffuse luminosity throughout the field. These phenomena are wholly analogous to those observed when a beam of light is passed through a dark room in the atmosphere of which fine dust particles are found. The path of the whole beam is made apparently uniformly luminous by the smaller particles, while occasionally there

appear points of bright illumination, due to the presence of larger particles. This is known as the Tyndall effect. The light which has passed through such fields is found to have become polarized.

It is evident that, in a solution whose particles are sufficiently large to become individually visible as points of light under the ultramicroscope, it immediately becomes possible to determine the size of the particles on the assumption that these are all of the same size. The procedure consists in determining the following quantities: (1) the total number of particles in a given volume by the usual blood-count method; (2) the weight of the dispersed substance in a given volume by a chemical or other analysis; (3) the density of the dispersed substance, which is usually taken as equal to that in the undispersed state. This undoubtedly introduces an error in the computation, since, in all probability, the density increases in the dispersed state, owing to increased compression by surface tension. This error is probably small unless the degree of dispersion is very great. By this method, particles in colloidal gold solutions have been observed and counted whose diameters were as small as 10^{-6} mm. This represents about the limits of individual visibility under the ultramicroscope, that is, with particles much smaller than this the field appears diffusely illuminated. This value is about one-hundredth that of the wave-length of violet light, and about ten times that of the calculated diameter of the ethyl alcohol molecule.

The rate of settlement under the influence of gravity has also been used to determine the size of colloid particles. By means of Stokes' law for the fall-rate of bodies through a viscous medium, a comparatively simple equation permits of the calculation of the diameter of the falling body when the fall-rate, the viscosity of the medium, and the densities, respectively, of the dispersed substance and of the medium are known. Perrin⁵ used the same principle in the preparation of suspensions in which the particles were all of more or less the same size, using, however, regulated centrifugation instead of simple settling under gravity.

There has also been developed, largely by Bechhod⁶ another method which throws some light on the relative sizes of particles, and also offers a very interesting and valuable experimental weapon for colloid investigation. This is the method of ultrafiltration. It has been found possible to produce graded filters which allow of the passage of particles below a certain size, and which restrain any larger ones. These filters are made by impregnating ordinary filters with gelatin and other colloidal solutions and drying with special precautions. The permeability decreases with the concentration of the gelatin or other substance used.

⁵ *C. R.*, 146, 967, 1908.

⁶ *Ztschr. Chem. Ind. Koll.*, 2, 3, 1907; *Die Kolloide in Biologie u. Medizin*, Dresden, 1912.

The investigations as to the size of particles all lead to two general conclusions: first, suspensions and colloidal solutions in general differ from one another mainly in degree of dispersion, at least up to the limit of individual detectibility of the particles under the ultramicroscope, beyond which point at present all is speculation, although the presumption is strong and the belief is growing that there is also no other fundamental distinction to be drawn between colloidal and so-called true solutions; second, it is always found that unless special purification is resorted to a colloidal solution contains particles of widely differing sizes side by side.

2. THE BROWNIAN MOVEMENT.—About a century ago the English botanist, Brown, noticed that very small spores and other bodies when suspended in water, and observed under the microscope, were in a state of rapid oscillatory and rotary motion. This motion of small masses of matter has come to be known as the Brownian movement. It is noticed in colloidal solutions whose particles are not too large, and at the same time are large enough to be individually detectible under the ultramicroscope. As a result of the theoretical considerations of Einstein,⁷ of Smoluchowski⁸ and of Corbino,⁹ and of the experimental researches of Svedberg¹⁰ and of Perrin,¹¹ the Brownian movement has come to be considered as nothing more nor less than a manifestation of that kinetic energy with which all matter is endowed, and which forms the basis of the kinetic theory of gases. A rapidly gyrating and oscillating colloid particle is therefore looked upon as a large scale picture of the state of the molecules themselves. These investigations have probably done more than anything save the development of the kinetic theory itself to place molecular and atomic speculations on a firm basis of plausibility.

Svedberg's investigations were instituted to determine the mean velocity of colloid particles whose mass could be determined by the ultramicroscopic method above referred to. Computing from these factors the average kinetic energy of the particles, this, according to Svedberg, gives the same value which would be computed for the particle on the basis of the kinetic theory. Perrin attacked the problem from a somewhat different point of view. The number of gas molecules in the atmosphere decreases from the surface of the earth outward at a rate which is determinable by computations based on the kinetic theory. Perrin set himself the task of determining the rate of decrease in the concentration of the particles of a colloid solution, in which the particles were of uniform size, the concentrations being determined at different levels in a cylinder in which the solu-

⁷ *Ann. der Phys.*, 9, 417; 11, 170; 17, 549; 19, 371.

⁸ *Ann. der Phys.*, 21, 756.

⁹ *Nuovo Cimento*, 20, 5.

¹⁰ *Ztschr. f. Elektrochem.*, 12, 853, 1906.

¹¹ *C. R.*, 146, 967, 1908.

tion had been allowed to stand until it had reached equilibrium with the gravitational forces. The result was that the same law of distribution was found to hold in this case as in the case of the atmosphere. The kinetic theory is thus shown to apply quantitatively as well as qualitatively to colloidal solutions.

3. ELECTRICAL PROPERTIES.—If a U-tube be filled with water, electrodes placed in each arm, and these electrodes maintained at a constant difference of potential either by a battery, dynamo, or other source of direct current, it is noticed that there is a continual flow of liquid in the tube, in one direction near the walls and in the opposite direction in the interior of the tube. There is every reason to believe that such currents will be set up in all cases whatever the nature of the liquid or of the tube, although the current set up may in particular cases be very small and even very rarely approach or equal zero. If we name the current along the walls simply the "current," and that through the interior the "countercurrent," then in the case of glass and water the direction of the current is from anode to cathode, and that of the countercurrent is from cathode to anode. This phenomenon is explained by the hypothesis that at the surface of contact between the glass and the water there is established a difference in electrical potential, the glass becoming negatively and the water positively electrified. If this assumption is valid it follows that if a particle of glass placed in water be subjected to the influence of two electrodes placed in the water, it will, being negatively charged, be attracted by the positively charged electrode (the anode) and repelled by the negatively charged electrode (the cathode). The result would be a wandering of the particle of glass through the solution toward the cathode. This result is confirmed by ample experiments. Furthermore, the phenomenon is common to all particles in all liquids, so far as is known, so that any colloidal solution placed in a potential gradient will show wandering of its particles in one direction or the other. Thus in water, ferric hydroxid, chromium hydroxid (and most hydroxids in the colloidal state), methyl violet, and some other dyes wander to the cathode. All colloidal metal solutions, sulphur, the halogen salts of silver, chlorophyll, rosin, mastic, most dyes, and, in fact, the great majority of substances investigated wander toward the anode. Albumin (and probably some other substances) wanders toward the cathode in acid solution, and toward the anode in alkaline solution. As will be seen later, the hypothesis of the existence of such electrical charges on colloid particles has been of very great use in explaining many forms of conduct on the part of dispersed systems.

4. SURFACE TENSION.—If a globule of mercury be divided into two parts, these two parts will unite again if opportunity be given. All the opportunity which is necessary, if the surfaces be clean, is to bring the two parts into mechanical contact. The union of the sep-

arate parts may, however, occur in a variety of other ways, in fact, in any way whatever whereby such union is physically possible. Thus, if the separate portions be of different size, the smaller one will have a higher vapor pressure than the larger, and evaporation from the smaller to the larger will take place until the whole of the smaller portion has transferred itself to the larger one, and the reunion is therefore complete. There is every reason to believe that if the two portions of unequal size were made electrodes in a galvanic cell, and this cell were then short circuited, that the smaller portion would go into solution and again deposit upon the larger one. In case the two portions were of the same size, these forms of recombination, with the exception of that of direct coalescence, would not occur if all other conditions were kept constant, but a slight difference of conditions in respect to the two portions would start the act of recombination, which would then in general proceed to completion. The same tendency is noticed with all substances. Thus in a liquid small crystals disappear while larger ones grow at their expense, and it may be stated that, other influences for the moment ignored, the most stable configuration which can be assumed by a given mass of any substance is that in which all of the substance is in one portion, and that portion is spherical in form. This is equivalent to saying that all bodies so arrange themselves as to expose the least possible surface. The force which tends to bring about this condition is called surface tension. In so far as surface tension alone is concerned it follows that any colloidal solution must be unstable, and tend to condense itself until all of the dispersed matter has aggregated itself together into a single mass of spherical form.

But there are many other forces which may under certain conditions act against surface tension. If the dispersed substance is one that is crystalline, the directive forces of crystallization overcome those of surface tension, and the form of stable configuration will be that of the crystal instead of spherical, and equilibrium will be established when all of the available substance has aggregated itself together into one large crystal. We know, on the other hand, a great many colloidal solutions which seem to be quite stable even in very high degrees of dispersion. To explain such cases we must look for other forces working against the force of surface tension. If the dispersed particles in a colloidal solution are all charged with the same kind of electricity, they will then repel one another with a force which will vary inversely as the squares of their distances from one another. This repulsion will then tend to work against any coalescence or other sort of union between the dispersed particles. We have already seen that colloidal particles are in general charged either positively or negatively, and this may be taken to some extent as explaining the stability of such systems. Equilibrium results when the surface tension is just counterbalanced by the electrical

repulsion. The extension of this idea has been of great value in colloid investigation. The electrical repulsion will not, of course, necessarily prevent the smaller particles from dissolving and depositing upon the larger ones, unless the solubility is affected by the charge. Concerning this we know nothing. The fact that colloid substances possess little or no solubility in the ordinary sense of the word means such solution and deposition must, of necessity, be a very slow process, and the colloid solution would thus appear to be perfectly stable over very long periods of time. There are many who believe that all such systems are only apparently stable, and that on account of the absence of any sufficiently rapid means of transformation which would allow the stabilizing influences to operate rapidly enough to be perceptible.

Chemical Properties of Colloids.—1. It is reasonable to suppose that the chemical properties of colloid solutions are very much what is to be expected from the chemical nature of the dispersed substance as it is known under other conditions. The colloidal solutions of arsenic sulphid should therefore react very much as would be expected of arsenic sulphid in general, except in so far as the substance is in a finely divided state in the presence of a dispersing medium (water) in which it is little soluble. Thus colloidal arsenic sulphid is soluble in alkalies and alkaline sulphid just as is the massive form. If a rod of zinc is suspended in a colloidal solution of arsenic sulphid there takes place a slow reaction, lasting over weeks and even months, whereby the sulphur of the sulphid unites with the zinc to form colloidal zinc sulphid, while a black deposit, probably arsenic, is found on the zinc. Chemical reactions with colloids are thus, as a rule, very slow, as is to be expected, but otherwise not essentially unusual.

2. The exact chemical composition of the disperse phase in a colloidal solution is probably not definitely known in any case. In the case of colloidal metal solutions, such as gold and silver, the suspended particles seem to be practically pure metals, but in most cases the composition is very problematical. The great variation in the properties of such solutions with variations in the methods of preparation are undoubtedly to a great extent due to small differences in composition. Thus the properties of arsenic sulphid vary greatly with the extent to which free hydrogen sulphid is removed from the solution, which is probably due to the differences in the amount of hydrogen sulphid absorbed or otherwise held by the arsenic sulphid. Linder and Picton believed that amorphous copper sulphid was a definite compound of copper sulphid with hydrogen sulphid. It has also been found that amorphous copper sulphid suspended in water continually deposits free sulphur, the cupric sulphid being at the same time largely converted to cuprous. It seems to be rarely or never the case that the disperse phase may be looked upon as a sub-

stance of definite composition, being usually, if not always, a more or less complex mixture of absorption products.

3. Although not usually pure substances, it is not at all un- plausible to assume that the dispersed particles may, to some extent, undergo ordinary electrochemical ionization, in which case the particles would partake of the nature of enormously large ions. This assumption is interesting as offering a purely electrochemical explanation of the origin of the charge which is found on such particles, and it is to be said that frequently the effect of foreign substances on the electrical charges of suspended particles is explainable on this assumption. For further information on these matters reference must be had to the papers of Duclaux,¹² Jordi,¹³ and P. P. von Weimarn.¹⁴

The Flocculation of Colloids by Electrolytes.—1. When neutral salts are added to colloid solutions in gradually increasing amounts there always follows sooner or later a precipitation of the dispersed substance. If the salt solution be removed and pure water added in its place, this decanted, and pure water again added, so as to wash out the salt as thoroughly as possible, the final result may be that the coagulum (or gel) becomes redispersed, or such redispersion may not occur. The outcome is determined both by the nature of the colloid and by that of the neutral salt used.

2. If acids and alkalies are the electrolytes used, the relationships are somewhat different. The addition of alkali to a negatively charged colloidal solution renders it more stable, while a positively charged colloid is flocculated. With acids the reverse of this condition holds; that is, the positive colloid is stabilized and the negative one is flocculated.

3. The concentration of the electrolyte required to flocculate a given colloidal solution depends very greatly on the nature of the electrolyte used. It has been generally considered that the cation of the electrolyte is the active agent in flocculating negative colloids, while the anion is active in the case of positive ones. Thus acids (hydrogen ions) are very effective in flocculating arsenic sulphid, and alkalis (hydroxyl ions) are effective with ferric hydroxid, as might be inferred from the preceding paragraph. Different cations, however, show very different degrees of precipitating or flocculating power. Thus in the case of arsenic sulphid, if the flocculating power of the potassium ion is taken as unity, that of calcium is about twenty, while that of the aluminium ion is three hundred and fifty. The flocculating power in this case increases very rapidly with the

¹² "Thesis," Paris, 1904.

¹³ *C. R.*, 136, 680, and 1,448; 137, 122; *Bull. Soc. Chim.*, 31, 573.

¹⁴ Articles in *Ztschr. Chem. Ind. Koll.*, 1906-1911. Also in "Grundzüge der dispersoid Chemie," Dresden, 1911.

valence of the ion, a relationship which seems to be quite generally true. It has been quite commonly considered that the effect of the anion in the flocculation of negative colloids is negligible. That this point of view is not tenable has recently been strikingly shown by Sven Oden¹⁵ in his work on colloidal sulphur. He finds that the effect of electrolytes on sulphur sols, which, like arsenic sulphid, are negative colloids, is distinctly the resultant of two factors, one a flocculating effect on the part of the cation, the other a dispersing effect on the part of the anion. It is not improbable that our views in regard to the phenomena of electrolyte flocculation will undergo considerable modification in the near future, as they are based on rather scanty experimental evidence. Since the properties of sols of the same materials differ very considerably with the most minute details of their method of preparation, it is naturally difficult for the same investigator even to obtain uniform results.

4. The actual concentration of a given electrolyte required to flocculate a sol depends also very greatly on the nature of the sol itself. Some sols are precipitated by very small concentrations of electrolytes (three to four one-hundredths normal acid being usually sufficient for arsenic sulphid), while gelatin, albumin, and protein substances in general require far higher concentrations. So marked is the difference in many cases that attempts have been made to classify colloids on the basis of their conduct in this respect. Thus colloids which are very sensitive to electrolytes are called *suspension colloids*, while those that are not very sensitive are called *emulsion colloids*. To the first class belong all the true, rather coarse-grained suspensions, while the sols that yield soft gelatinous flocculates usually fall into the second class. It is also very frequently true that the latter type shows the phenomenon of reversible flocculation (see *ante*). This classification is for many purposes quite useful, but cannot be considered as very fundamental. For example, if the electrolyte used be a salt of a heavy metal most of the so-called emulsion colloids, such as albumin, are irreversibly flocculated.

5. The flocculation of sols by electrolytes is usually explained as due to the phenomenon of absorption. That is, the flocculating ion is absorbed from the solution by the dispersed particles. Since in general the ion which is absorbed is the one whose electrical charge is opposite to that of the dispersed particle the absorption results in a reduction of the charge on the particle, and allows the aggregating forces of surface tension to become operative. The evidence of the validity of this assumption is considerable. Thus the flocculated colloids always contain appreciable amounts of the ion which caused the precipitation, which is *prima facie* evidence of the absorption. Furthermore, the electrical charges on the particles may be measured by determining their rates of migration, and the effect of elec-

¹⁵ "Inaug. Diss.," Upsala, 1913, pp. 118 *et seq.*

trolytes on this charge may also be observed. It is very commonly true that the addition of a precipitating electrolyte to a sol reduces materially the charge of the particles. This is not, however, always true, and the relationships are more complicated than has generally been assumed. Nor is it always true that the complete neutralization of the electrical charge on dispersed particles results in flocculation. For example, acetic acid may be added to arsenic sulphid sols in such quantities as to completely neutralize the negative charges of the particles and, further, so much acid may be added that the particles acquire a very considerable positive charge, all this without the least signs of flocculation. Ultimately so much acid may be added as to cause flocculation.

6. In the flocculation of sols by electrolytes there is frequently observed a curious effect known as the "*zone-phenomenon*." It is observed when increasing amounts of certain electrolytes are added that at a certain concentration flocculation will be brought about, while if the concentration be greater flocculation will not occur, although still further increase of concentration will result in another flocculation zone. The phenomenon is most common when the electrolyte used is a salt that shows marked hydrolysis, such, for example, as ferric chlorid. If a negative sol be treated with a solution of this salt it is obvious that there will be three precipitating influences present, namely, the hydrogen ions and the colloidal ferric hydroxid, both of which are formed by the hydrolysis of the ferric chlorid and the unhydrolyzed ferric chlorid. Since the amounts of these precipitating substances in a given solution vary with the concentration, and since each has its own concentration function in precipitation, it will be seen that the zone phenomenon may be accounted for in such cases. Since, however, the zone phenomenon occurs in many cases where strongly hydrolyzed electrolytes are not used, as, for example, in the agglutination of bacteria by citric and some other acids, the explanation is not wholly sufficient. There are also many curious phenomena concerned with the action of electrolytes on sols which have as yet been very little investigated, and which will probably throw considerable light on the subject.

The Mutual Reactions of Colloids.—The conduct of mixtures of two different sols is of very great interest and variety, both in the absence and in the presence of electrolytes. A number of particular cases may be distinguished, and these will be taken up seriatim.

1. When two positive or two negative sols are mixed together nothing very much seems to happen. It is generally considered that the addition of one sol to another of like electrical properties results in no action. Whether this is wholly true or not is doubtful, but it is at least true that in all cases investigated up to the present time

neither individual nor mutual flocculation occurs. Whether the presence side by side of two sorts of similarly charged disperse particles results in any change in the dispersion of either is not known. Except to show that no mutual precipitation occurs, these cases have been but little studied.

2. When two oppositely electrical colloids are mixed mutual precipitation may or may not occur. The factor which in the main determines the outcome is the relationship between the amounts of the two colloids used. If neither is present in too great excess complete mutual precipitation in general occurs. If either is present in great excess precipitation does not in general occur.

3. The effect of a great excess of one colloid in preventing mutual precipitation is very marked when the colloid in excess is one of the emulsion-colloid type. Thus gelatin, a typical emulsion colloid, when present in excess over another colloid very frequently prevents all flocculation, even in fairly coarse-grained suspensions. Advantage has been taken of this action in preventing scaling in boilers. This scaling is due largely to the fact that lime salts held in solution as bicarbonates are decomposed by heat, with the separation of calcium carbonate, at first in the highly dispersed state. This gradually aggregates together and deposits on the interior of the boiler as an amorphous flocculated colloid, which in time becomes very compact, and in many cases crystalline. If, however, a small amount of glue (impure gelatin) be added to the boiler water the colloidal constituents of the water do not flocculate and compact, but remain suspended, and may from time to time be blown off. Another illustration is found in the preparations of photographic emulsions. The silver halides flocculate very readily in pure water, but in gelatin solution remain in a highly dispersed state, which is necessary to the preparation of the plate. In this case, not only is the suspension protected from flocculation, but also a degree of dispersion is reached which is far beyond anything attainable in pure water. The same is true when lysalbinic acid is used to prevent flocculation of colloidal silver. In pure water only very dilute suspensions of metallic silver are obtainable, but in the presence of lysalbinic acid suspensions containing as high as ninety per cent. of silver are obtainable, the product being used medicinally under the name of "argyrol." These are the phenomena known as "*protective actions*," and the gelatin, albumin, or other colloid which exerts the protective action is spoken of as a "*protective colloid*."

4. The protective action of certain colloids is not only exerted against the tendency of the protected colloid to spontaneously flocculate, but also a certain and very great protection is offered against flocculation by electrolytes. Thus Zsigmondy¹⁶ was able to find a definite measure of the protective action of certain colloids on the

¹⁶ *Ztschr. f. analyt. Ch.*, 40, 697, 1902.

precipitation of gold suspensions by means of sodium chlorid. The method was to find the amount of the protective colloid that was just necessary to prevent the flocculation of a fixed amount of a given gold sol by a fixed amount of the salt. In this way it was observed that the protective action of different colloids is very different. Thus if the amount of starch in solution which is necessary for protection be taken as 2,500 the amounts of various other colloids required are as shown in the following table:

| | | | | | | |
|------------------------|--------|---------|------------|---------|---------|------|
| Protective colloid.... | Starch | Dextrin | Gum Arabic | Albumin | Gelatin | Glue |
| Amount required | 2,500 | 1,200 | 40 | 25 | 1 | 1 |

5. Very simple theories are devisable to explain the interactions between different colloidal solutions. Thus two oppositely electrical colloids may be considered to precipitate one another mutually because of the electrical attraction existing between all oppositely charged particles. This results in bringing together the oppositely charged particles with the formation of relatively neutral aggregates, which, as shown in the discussion of the flocculation by electrolytes, is a condition favorable to precipitation. For very obvious reasons, then, no flocculation would be expected when two like charged colloids are mixed. Many objections may be made to the unqualified acceptance of this explanation. It offers, nevertheless, a valuable leading idea when not accepted too dogmatically.

A number of factors probably contribute to the protective action of many colloids. To some extent the effect may be purely mechanical, since increased viscosity imparted by the presence of the protective emulsion colloid will shorten the mean free path of the flocculable particles, and thus materially lessen the probability of impacts between them. Consequently flocculation will not occur as readily. Further, the ultramicroscope gives considerable evidence of the existence of another and very important factor. It seems certain, at least in many cases, that the protective colloid arranges itself in a film or coating around the flocculable particles, and in this way prevents the aggregation of the particles. These factors are not, however, sufficient to explain all cases of protective action. It must be considered that in some cases, at least, the protective colloid exerts a truly dispersive effect, such as would result from a nullification of surface tension forces. The ease with which a small amount of soap will emulsify a large amount of oil is difficult to explain on any other hypothesis. When an oil is shaken with pure water little or no emulsification results, while in the presence of the soap as a protective colloid the same amount of work in shaking accomplishes enormously greater results. In fact, the oil will spontaneously emulsify by merely standing in contact with the soap solution. The equilibrium condition of oil in contact with pure water is reached

when the oil is very little, if any, dispersed, while in contact with soap solution equilibrium is reached only when the oil is highly dispersed. From the surface tension point of view this would be expressed by saying that in contact with pure water the surface tension is large and positive, while in contact with soap solution the surface tension is large but negative. It is impossible to say to what extent these effects may be due to obscure chemical action.

The Preparation of Colloidal Solutions.—1. This subject might perhaps upon logical grounds have best been treated in an opening paragraph. However, with the conclusions that have been reached in the foregoing discussion the whole matter may be dismissed very briefly. The conditions which must be fulfilled in order to obtain colloidal solutions are in the main summarized in the following paragraphs:

2. A medium must be chosen in which the given substance does not reach to any great extent the maximum, molecular degree of dispersion—that is, in which what is usually called true solution does not occur to any great extent. While, as pointed out at the beginning of this discussion, there is no sharp line to be drawn between colloidal and true solutions, the substances that are distinctly recognizable at the present time as colloidal solutions carry particles which are in the neighborhood of one thousand times as large as average molecules. From media in which the dispersion is approximately molecular the dispersed substance usually shows a strong tendency to separate in the crystalline form, although it may frequently, if not generally, first appear in the colloidal form, then more or less rapidly becoming crystalline. The only distinction to be drawn is this: from solutions in which the dispersion is very great, approximating the molecular, separation in short time in the crystalline form is favored; from solutions in which the dispersion does not approximate the molecular separation persistence in the amorphous state is favored.

3. A colloidal sol or gel, one or the other, may generally be produced from any substance in any medium in which the amount of molecular dispersion is at a minimum by means of any reaction whereby the new substance is produced from solution. Thus, for example, arsenic sulphid does not disperse in water to molecular extent in any considerable degree. Therefore, by mixing together a solution containing an arsenic compound which is soluble, and a solution of hydrogen sulphid, the resulting arsenic sulphid will appear in the colloidal state. Whether it appears in the dispersed state as a sol or in the flocculated state as a gel will depend mainly on the electrolyte content of the solutions which are used. If a solution of arsenic chlorid be used the resulting solution will contain considerable free hydrochloric acid, and the tendency will be for the

resultant arsenic sulphid to appear in the flocculated or gel form. If it is desired to prepare the substance in the sol form electrolytes must be avoided. This may be done by using a solution of arsenic trioxid instead of one of arsenic chlorid.

Any reaction which is brought about under the above conditions will result in the formation of a colloidal product. The dialysis of salts which form insoluble hydroxids simply allows the normal reaction of hydrolysis to complete itself. The resulting hydroxids appear in such a way as to fulfill the above conditions, and consequently appear in the colloidal state. In this connection note the preparation of colloidal sodium chlorid (see *ante*).

4. In an appropriate medium most if not all substances, crystalline or otherwise, may be brought directly, that is, without the intervention of specific chemical reactions, into the colloidal state. In some cases this may be accomplished by mechanical means. Thus oils violently shaken with water disperse to some extent and form emulsions of greater or less stability. By shaking glass, quartz, and the like with various liquids in which they are virtually insoluble the abrasion results in the formation of more or less dispersed systems, usually not very stable.

Many metals may be brought into the dispersed state by causing an electric arc to pass between points held under a liquid. This electrical dispersion method has been very considerably used, but is obviously confined to substances which are conductors of electricity.

On the other hand, many substances when merely brought into contact with an appropriate medium spontaneously undergo dispersion. Thus gelatin, glue, tannin, and many other substances spontaneously disperse in water. Even crystalline substances frequently do this. Thus soaps which have been crystallized from alcohol solutions when brought into water disperse in the colloidal state. Crystallized cuprous sulphid, the mineral known as chalcocyste, disperses in the colloidal form in solutions of hydrogen sulphid.

Substances which go spontaneously into the dispersed colloidal state are usually spoken of as "*lyophillic*," while those that tend to spontaneously leave the dispersed state are called "*lyophobic*." It is evident that a substance may be lyophillic with respect to one medium and lyophobic with respect to another. Furthermore, a very small change in the nature of the medium may cause the change from a lyophillic to a lyophobic colloid. Thus oils are lyophobic with respect to water, but lyophillic with respect to even very dilute soap solutions.

Applications to Biology.—1. When one considers the relatively infrequent occurrence in biological systems of either crystalline substances, or of substances that may be readily made to crystallize from water (the universal biological dispersing medium), it immediately becomes evident that the chemistry and physics of such

systems must be in the main colloidal. All biochemistry is thus in the main colloid chemistry. Aside from mineral salts, urea, uric acid, and a few other bodies, the reagents and products which are active in life processes are all to be found in the living organism in the colloidal state. While crystalline directive forces are in general absent, we have nevertheless to deal in biological phenomena with a great variety of directive forces of a wholly different character. Colloidal substances in high degrees of dispersion such as proteins and the like in the alimentary fluids are being continually converted into active living tissues, a process manifestly involving very definite directive forces, since the product (the tissue cells) is an organized one, even though its organization is not similar to that of a crystal. Thus the building of living tissue involves among other things the conversion of sols of many sorts (alimentary fluids, blood, etc.) into gels. In other words, the living tissue is to a certain extent to be looked upon as a colloidal gel, differing, however, from laboratory gels in possessing a definitely organized cell structure. While it is true that in the spontaneous gel formation with certain colloids, as, for example, gelatin, myelin, or web structures are formed purely as a result of the physical and chemical forces active, these cannot be said to bear any very strong resemblance to living cells. It is thus apparent that life processes differ very materially from those of the chemical laboratory. On the other hand, it is true that many of the component reactions which go to make up the life process may be very closely duplicated by laboratory means, and that already the study from the colloid chemistry point of view of the reaction of many of the substances which go to make up the living organism has given interesting and important results. Furthermore, the whole field of colloid investigation has been greatly stimulated by the hearty support and encouragement which it has received from biologists. Some slight attempt will be made here to illustrate by a few examples the far-reaching possibility of explanation which colloid chemistry offers of certain phases of biological science. The actual accomplishment in the field is already so great that only a very limited discussion can be offered here.

2. The action of electrolytes on emulsions of bacteria is wholly analogous to their action on colloidal suspensions. The bacterial emulsions are very sensitive to flocculation by mineral acids, one thousandth normal hydrochloric and sulphuric acids usually being sufficient to cause complete clumping and settling of the bacteria. Neutral salts, with the exception of those of silver, mercury, iron, and aluminium, do not flocculate the bacteria. If, however, the bacteria are first treated in the absence of electrolytes with an agglutinating serum small concentrations of salt solutions will bring about flocculation. It is also known that bacteria have the power of absorbing

agglutinins from sera, so that it is evident that what we have here is a case of the production of a flocculable combination of bacteria and agglutinin, neither component of which is alone flocculable.

Citric acid in concentrations ranging between one one-hundredth and one eight-hundredth normal produces flocculation. In either greater or less concentrations no flocculation is produced, which is an illustration of the so-called zone phenomenon.

Like all other suspensions, bacteria are electrically charged, and consequently wander in the electric field. Under all ordinary conditions the charge which they carry is negative, from which in their general conduct it might be expected that they would conduct themselves similarly to arsenic sulphid, which is also negatively charged. This is found to be the case. Their rate of migration is reduced by acids and evidently somewhat increased by alkalies, although very little alkali is necessary to bring about disintegration of the bacteria.

It has also been found that all colloids are more or less sensitive to light in respect to their migration rates. Bacteria also show this phenomenon, as they migrate notably slower in the light than in the dark. It is quite possible that this reduction in their electrical charge may to some extent be responsible for the bactericidal action of light.

3. A very interesting application of the principles of colloidal precipitation by electrolytes has recently been made by Loeb.¹⁷ He finds that the eggs of the *Fundulus*, a small fish, are killed by being immersed in a pure isotonic salt solution, in spite of the fact that they normally develop in sea water. The factor which allows of their development must therefore be sought in some other constituent of the sea water which is absent in the pure salt solution. This Loeb finds in the presence of small amounts of calcium salts. Further, if a small amount of calcium chlorid be added to the pure salt solution the eggs will develop in it as well as in sea water. Loeb's explanation is simple and very ingenious. He supposes that the sodium chlorid is toxic, provided it can diffuse into the egg. In the absence of calcium salts such diffusion is possible because the sodium chlorid is not a sufficiently powerful colloid precipitant to make the membrane about the egg impervious. It is, however, very well known that calcium salts (and all bivalent cations) are far more effective colloid precipitants than sodium ions. Consequently the presence of a relatively small amount of calcium chlorid in the salt solution is sufficient to so condense the egg membrane as to make it impervious to the sodium chlorid, and thus render the latter non-toxic.

4. Another interesting application of colloidal principles is found in what is known as the Danysz phenomenon. It is found that the neutralization of the toxicity of diphtheria toxin by the

¹⁷ *Am. J. Physiol.*, 6, 411, 1902.

antitoxin depends on the way in which the two are mixed. If a quantity of toxin just sufficient to neutralize a fixed amount of antitoxin when it is added all at once be in another experiment added in small installments, the resulting mixture will be found to be still quite strongly toxic. This is quite analogous to what is found in the interaction of many colloids. The amount of a given colloid required to neutralize and precipitate another depends greatly on the way in which it is added.

5. Römer,¹⁸ Field and Teague,¹⁹ and Teague and Buxton²⁰ all carried out interesting investigations directed toward determining the migration directions (electrical charges of toxins and antitoxins). Their conclusions were that all wandered toward the cathode, and that all were therefore positively charged. If this is correct the analogy between toxin and antitoxin reactions and those of simple colloids is rather mutilated, since two positive colloids are not supposed to react with one another. It is, however, more than possible that the above experiments are misleading. In all cases agar diaphragms were used. Through these there would always occur a streaming of water toward the cathode as a result of the electrical potential between the agar and the water. This might well be so great as to obscure, and even more than overcome any anodic wandering that might occur. Furthermore, the conduct of proteins in general in the electric field is a very complex one, and one that is only just beginning to be understood. For these reasons it is at present very dangerous to draw any very dogmatic conclusions.

6. In closing mention may be made of what seems to be an immunity phenomenon which seems rather clearly to be a case of protective colloid action. It is observed when an agglutinin is added to a bacterial emulsion that if an excess of the agglutinin be added no agglutination occurs. This is wholly analogous to the fact that, while a small amount of gelatin will precipitate arsenic sulphid suspension, a larger amount will not.

Conclusions.—While great progress has been made in the field of colloid investigation from the chemical and physical sides, and while also many very striking analogies are to be found from the biological side, it is nevertheless true that we are still very much in the dark in regard to a great many matters. The one great difficulty which lies in all such investigations is that it is a matter of very great difficulty to duplicate results. The nature of any colloid sol or gel depends so greatly upon its whole previous history, apparently down to the least detail, that great discrepancies in experimental results are found. Even the age of a sol is frequently a matter of very great importance in determining its properties. For example

¹⁸ *Berl. klin. Wochenschr.*, Vol. 41, p. 209, 1904.

¹⁹ *Journ. Exp. Med.*, Vol. 9, pp. 86 and 223.

²⁰ *Ibid.*, Vol. 9, p. 254.

a freshly prepared gelatin solution will not precipitate arsenic sulphid, but it will do so after it has stood for some hours. What is now greatly needed is more data on a greater variety of colloids that have heretofore been investigated and work directed toward the preparation of colloidal solutions of definite character. Until something has been accomplished in these directions all biological analogies and the like cannot be anything more than qualitative, and the same holds true for many of the physical and chemical conclusions which have been discussed in this chapter.

INDEX OF AUTHORS

- Abderhalden, 95, 98, 493-496
 Abel and Ford, 96
 Abramow, 41
 Abt, 387, 394
 Adami, 24, 134, 234, 281, 282
 Addis, 171, 303
 Adler, 388
 Admiradzibi, 411, 432
 Altmann, 184
 Amoss, 436
 Anderson, 362, 365, 368, 373, 374, 376-382, 389, 390, 401, 411, 426, 428, 430, 437, 463, 464
 Anderson and Goldberger, 54
 Anderson and Schultz, 365
 Andrejew, 372, 437
 Apolant, 373
 Arima, 34, 477
 Aronson, 473
 Arrhenius, 120, 122
 Arrhenius and Madsen, 120, 121, 122
 Arthus, 361, 368, 404
 Arthus and Breton, 361
 Asakawa, 46
 Aschoff, 127
 Ascoli, 148, 237, 268, 493
 Ascoli and Izar, 496, 497
 Auer and Lewis, 364, 390
 Axamit and Tsuda, 319
- Bab, 210
 Babes, 65, 440
 Babes and Broca, 439
 Babes and Lepp, 74
 Bach and Chodat, 184
 Baecher, 316
 Baginsky, 473
 Bail, 5, 11, 20, 21, 80, 228, 229, 289, 326, 443
 Bail and Kleinhans, 76
 Baldwin, 442
 Bandeller and Roepke, 351, 356, 357
 Bang, Ivar, 44, 47, 195
 Bang and Forsmann, 97
 Banzhaf, 430
 Banzhaf and Famulener, 379
 Banzhaf and Steinhardt, 379
 Barber, 15, 67
 Bartel, 13
 Bartel and Neumann, 343
 Bauer, 209, 442
 Baumann, 83
 Baumgarten, 84, 135, 136
 Bechold, 185, 242, 243, 266, 504
 Beck, 53, 439
 von Behring, 65, 73, 82, 83, 84, 85, 104, 107, 359, 390, 407, 458, 459, 472
- von Behring and Kitasato, 73, 75, 86
 von Behring and Kitashima, 407
 von Behring and Wernicke, 66, 73, 75, 86
 Belfanti, 5
 Belfanti and Carbone, 91
 Beniasch, 246
 Bergel, 204
 Berghaus, 447
 Bertin, 426
 Bertrand, 75, 86, 105, 464
 Besredka, 46, 67, 71, 75, 132, 133, 198, 349, 374, 375, 378-380, 387-390, 401, 431, 432, 476, 484, 487
 Besredka and Steinhardt, 368, 374, 377
 Bessau, 402, 476
 Besson, 289
 Bickel, 323
 Biedl and Kraus, 365, 368, 369, 383, 390, 402, 404
 Biggs, 324, 449-451
 Billitzer, 266
 Billroth, 135
 Biltz, 123, 242
 Bispham, 319, 333
 Bizzozero, 234
 Blackstein, 8
 Blair, 444
 Blumenthal, 131
 Boas, 200, 210, 211
 Boehme, 139
 Bogomolez, 371
 Böhm, 44
 Bolton, 92
 Borden, J. H., 223
 Bordet, 89, 91, 94, 122, 123, 140-145, 153, 154, 156, 158-160, 164, 165, 170, 186, 223, 239-243, 245, 248, 288, 296-298, 311, 416, 473
 Bordet and Gay, 166, 167
 Bordet and Gengou, 186, 188
 Bordet and Streng, 167
 Borrell, 280, 478
 Bouchard, 20, 85
 Boycott and Douglas, 239
 Boyle, Robert, 1
 Bradley, 341
 Brand, 179, 180
 Brau and Denier, 34, 87
 Braun, 200, 205, 411
 Breton, 361
 Brieger, 29, 30, 77, 337, 338, 404
 Brieger and Fraenkel, 73
 Brieger and Mayer, 70
 Briscoe, 278, 279
 British Plague Committee, 479
 Broca, 439
 Brodie and Dixon, 369
 Bronfenbrenner and Noguchi, 181, 183
- Brown and Fraser, 43
 Brown-Séquard, 405
 Browning, 155, 167, 177
 Browning and Cruikshank, 201
 Browning and McKenzie, 196
 Bruck, 192, 198, 199, 205, 439, 440
 Bruschetini, 41
 Buchner, 33, 80, 104, 125, 134, 137, 143, 288, 300
 Buchner and Hahn, 72
 Buchner and Orthenberger, 178
 Bujwid, 430
 Buller, 286
 Bullock, 341
 Bullock and Western, 318
 Bürgers, 39
 Bütschli, 294
 Buxton, 518
- Cagniard-Latour, 1
 Calmette, 75, 86, 87, 105, 106, 174, 356, 440, 464-466, 478
 Calvary, 369
 Canfora, 5
 Cantacuzene, 299
 Carbone, 91
 Carey, 183, 278, 279, 284, 306, 307, 343
 Carrière, 39
 Carroll, 55
 Castellani, 100, 232
 Casuto, L., 503
 Cattani, 84
 Chamberland and Roux, 66, 73
 Chantemesse, 438, 469, 475, 476
 Chapin, 318, 319, 320, 323
 Charrin and Roger, 89, 218
 Chauveau, 57, 66, 83
 Cherry, 104, 105
 Chirolanza, 8
 Chodat, 184
 Choksy, 479
 Christian and Rosenblatt, 442
 Citron, 21, 101, 198, 210, 440, 469
 Clappole, S., 281
 Clowes, 214
 Coca, 175, 398, 405, 406, 465
 Cohn, 77, 118, 145, 176, 182
 Cohnheim, Otto, 258
 Cole, 228, 468, 475
 Coic, Dochez and Gillespie, 221
 Cole and Meakins, 341
 Collins, 457
 Conradi and Drigalski, 219
 Conte, 13
 Contejean, 99
 Corbino, 505

- Cornet and Kossel, 60
 Courmont and Doyen, 131
 Cowie and Chapin, 318,
 319, 320, 323
 Cox, 324, 353
 Craig and Nichols, 203
 Craw, 124
 Crile, 398
 Cruikshank, 201
 Cumming, 367
 Currie, 428
 Curschmann, 60
- Dale, 398
 Danysz, 18, 123, 124
 Dautwitz, 97
 Dean, 190, 193, 316-318,
 321, 323, 480
 Delanôe, 411
 Delezenne, 92
 Denier, 34, 87
 Denys, 80, 90, 312, 326,
 351, 357, 473
 Denys and Havet, 168,
 300
 Denys and Kaisin, 300,
 308
 Denys and Leclef, 311,
 312
 Denys and Marchand,
 325
 Denys and Van der Velde,
 73, 87
 Descatello and Sturli, 237
 Detre, 199
 Deutsch, 100, 301
 Deutsch and Feistmantel,
 326
 Dick, 303
 Dickson, 44, 276
 Dieudonné, 447, 487
 Dineur, 225
 Ditman, 341
 Dixon, 369
 Dochez, 221
 Dochez and Gillespie, 475
 Doelter, 501
 Doerr, 34, 87, 263, 372,
 380, 410, 422, 432,
 436, 469
 Doerr and Russ, 381, 382,
 388, 389, 391, 394,
 396, 400
 Doflein, 6
 Dold, 413, 415, 417, 418,
 421, 423
 Donath, 147
 Donath and Landsteiner,
 169
 Dönitz, 46, 130
 Döring, 196
 Douglas, 239, 313-316,
 334-337, 340
 Doyen, 131
 Draper and Handford, 54
 Dreyer and Madsen, 113
 Dreyfus, 375, 428
 Drigalski, 219
 Duclaux, 509
 Dunbar, 434, 435
 von Dungern, 92, 124, 143,
 145, 171, 195, 214,
 215, 263, 268, 269,
 429
 von Dungern and Coca,
 175, 465
 von Dungern and Hirsch-
 feld, 58, 239, 372, 436
 Durham, 89, 218-220, 229-
 231, 248
 Dwyer, 22, 228, 310, 402,
 476
- Eggers, 318
 Ehrlich, 37, 56, 57, 58, 75,
 83, 85-87, 93, 95, 106-
 120, 122, 124-126, 128-
 133, 141-144, 146, 147,
 149, 151, 152, 155,
 156, 158-160, 162, 164,
 173, 177, 182, 186,
 187, 189, 234, 235,
 239, 264, 301, 416,
 439, 440, 443
 Ehrlich and Bordet, 164
 Ehrlich and Brieger, 337,
 338
 Ehrlich and Marshall, 157
 Ehrlich and Morgenroth,
 142, 143, 146, 147,
 150, 151-155, 157, 165,
 177, 181
 Ehrlich and Sachs, 153,
 155, 165, 166
 Einstein, 505
 Eisenberg, 19, 229, 233,
 268, 429
 Eisenberg and Volk, 226,
 233, 235, 236
 Eisenbrey, 368, 369, 373,
 398
 von Eisler, 47, 132, 133,
 195, 196
 Elschnaig, 437
 Emery, 341
 Emmerich, 85
 Engelmann, 287
 Epstein, 58
 Epstein and Ottenberg, 239
- Falloise, 171, 303
 Famulener, 379
 Fassin, Louise, 172
 Faust, 30
 Ferran, 66, 345, 351, 484
 Ferrata, 178, 183
 Ficker, 220, 223
 Field, 513
 Fildes, 210
 Finsterer, 445
 Fischer, 7, 128, 135
 Fish, 94
 Fleischmann, 99
 Fleming, 324, 340
 Flexner, 359, 370
 Flexner and Jobling, 470
 Flexner and Noguchi, 174,
 465
 Flexner and Sweet, 45
 Flugge, 84, 87, 134
 Foa, 85
 von Fodor, 79, 134
 Ford, 96, 234
 Fornet and Müller, 257,
 261, 262, 267, 268
 Forsmann, 97
 Fraenkel, 24, 47, 73, 184
 Frank, 92
 Fraser, 43
 Freeman, 340
 Friedberger, 23, 38, 172,
 174, 190-192, 194,
 240-246, 270, 291, 366,
 378, 385, 391, 396,
 397, 399-402, 411, 413,
 414, 419-423, 441, 442
 Friedberger and Hartoch,
 394, 395
 Friedberger and Ito, 422
 Friedberger and Mita, 366,
 367, 415, 430, 432
 Friedberger and Moreschi,
 69
 Friedberger and Nathan,
 418
- Friedberger and Salecker,
 423
 Friedberger and Szmanow-
 ski, 418
 Friedemann, 148, 242-244,
 250, 265, 380-386, 390,
 395, 397, 399-401, 406-
 408, 442
 Friedemann and Isaak,
 403
 Friedemann and Sachs, 176
 Frosch, 5
 Futaki, 19, 80, 325, 326
- Gabritchewsky, 291, 311
 Gaffky, 487
 Galleotti, 70
 Galtier, 13
 Garbat and Meyer, 477
 Garre, 83
 Garrey, 292
 Gautier, 29
 Gay, 54, 68, 163, 166, 167,
 189-193, 212, 438, 484
 Gay and Adler, 388
 Gay and Claypole, 8, 291
 Gay and Rusk, 268, 269
 Gay and Southard, 364,
 371, 380, 382, 387-
 390, 394
 Gengou, 171, 172, 186, 188,
 190, 192, 211, 265,
 303, 304, 400, 493
 German Plague Commis-
 sion, 13, 479
- Gibier, 51
 Gibson, 430, 457
 Gibson and Collins, 457
 Gilbert, 7, 348
 Gildersleeve, 92
 Gillespie, 221, 475
 Glynn, 353
 Glynn and Cox, 324
 Goldberger, 54
 Gonzales, 430
 Goodall, 428
 Gottlieb, 30, 39, 40, 44,
 45, 99, 409
 Grafe and Graham, 148
 Graham, 148, 499, 502
 Gramenitski, 184, 185
 Grassberger, 458
 Grassberger and Schatten-
 froh, 34, 36, 73, 87
 Griffiths, 29
 Grohman, 79, 134
 Gruber, 80, 224, 306, 312,
 326
 Gruber and Durham, 89,
 218-220, 229, 248
 Gruber and Futaki, 19,
 325, 326
 Gruber and Wiener, 88
 Grünbaum, 220
 Guggenheim, 180
 Gumaleia, 52
 Gurd, 303
 Gutstadt, 439
- Haccius, 488
 Hada and Rosenthal, 148
 Haendel, 183, 194, 221,
 263, 351, 373, 395,
 406, 407, 474, 475
 Haffkine, 485, 486
 Hahn, 55, 72, 168, 300,
 304, 448, 460, 462, 480
 Hall, 463
 Hallier, 77
 Hamberger and Moro, 391
 Hammarsten, 28, 404

- Handford, 54
 Hankin, 168, 301
 Hardy, 242, 301
 Harriehausen and Wirth, 462
 Harris, 96
 Harrison, 225
 Hartoch, 394, 395
 Havet, 168, 300
 Hayem, 272
 Hecker, 175, 180
 Heidenheim, 369
 Heilner, 493
 Hektoen, 238, 280, 317, 319, 321, 322, 323, 325, 333
 Hektoen and Ruediger, 178, 314, 318, 395
 Helme, 276
 Henneguy, 275
 Hérincourt, 74
 Herman, 171, 303, 478
 Herz, 477
 Hewlett, 171, 303
 Hirschfeld, 58, 239, 372, 436
 Hiss, 218, 222, 230, 309, 310
 Hiss and Zinsser, 309
 Höber, R., 44
 Hodenpyle, 69
 Högyes, 66
 Holobut, 411, 432
 Hopkins, 239, 353
 Hopkins and Zimmermann, 202
 Hort, 344
 Hüne, 316, 317, 321
 Hunt, Reid, 409
 Hunter, John, 62, 79, 134

 Inmann, 316, 323
 Isaak, 403
 Isaëff, 85, 87-89, 137
 Isaëff and Ivanoff, 218
 Ito, 422
 Ivanoff, 218
 Izar, 415, 496, 497

 Jacobæus, 210
 Jacobsthal, 204
 Jacoby, 96, 250
 Jacoby and Schütze, 185
 Jaffe, 246
 Jagic, 265
 Jameson, Eloise, 242
 Jenner, 62, 63, 345, 481
 Jennings, 286
 Joachim, 227
 Jobling, 332, 470, 471
 Jobling and Peterson, 424
 Jochmann, 306, 307, 469, 470
 Jochmann and Müller, 87
 Joest, 73
 Johannesen, 426
 Johnston, 8, 193
 Joos, 69, 226, 240, 259
 Jordi, 509
 Jorgensen and Madsen, 338

 Kainsin, 300, 308
 Kaliski, 148, 149
 Kanthack, 289
 Kanthack and Hardy, 301
 Kantorowicz, 307
 Karasawa and Schick, 448
 Karlinski, 73
 Karsner, 373
 Kempner, 34, 73, 87
 Kempner and Pollack, 41, 131
 Kempner and Shepilewsky, 131
 Keysser and Wassermann, 422-424
 King, 375
 Kiss, 176
 Kisskalt, 20
 Kitasato, 32, 57, 73, 75, 77, 84-86, 104
 Kitashima, 407
 Klausner, 204
 Klebs, 272
 Klein, 239
 Kleinbans, 76
 Klien, 332
 Knaff-Lenz, 176
 Knorr, 46, 125, 407
 Robert and Stillmarck, 106
 Koch, 72, 76, 77, 296, 345, 355, 356, 439, 440
 Koehlich, 342
 Kohn, 443
 Kolle, 13, 56, 69, 83, 351, 482, 486
 Kolle and Martini, 479
 Kolle and Otto, 487
 Kolle and Schürmann, 39
 Kolle and Strong, 487
 Kolle and Wassermann, 469, 470
 Korschun and Morgenroth, 169, 306
 Kossel, 60, 94
 Kraus, 60, 70, 87, 90, 248-251, 365, 368, 369, 383, 390, 402, 404, 415, 421, 422, 469, 488
 Kraus and Admiradzibi, 411, 432
 Kraus and Doerr, 34, 87, 380, 410, 422, 432, 469
 Kraus, Doerr and Sohna, 263, 372, 436
 Kraus and Joachim, 227
 Kraus and von Pirquet, 264-266
 Kraus and Stenitzer, 477
 Kraus and Volk, 200, 389
 Kretz, 390, 408
 Krumwiede, 52
 Kruse, 20, 39
 Kumagai, 402
 Kyes, 174, 175, 465
 Kyes and Sachs, 174

 Lagrifoul, 477
 Lamar, 319, 332, 333
 Lambert, Adrian, 310
 Lambert, R., 277
 Lambotte, 171, 303
 Landois, 91, 405
 Landsteiner, 47, 92, 97, 122, 169, 195, 200, 238, 250, 265, 371
 Landsteiner and Dautwitz, 97
 Landsteiner and Donath, 147
 Landsteiner and von Eisler, 47, 132, 133, 195, 196
 Landsteiner and Jagic, 265
 Landsteiner and Levaditi, 54
 Landsteiner, Müller and Pötzl, 200
 Landsteiner and Richter, 237
 Landsteiner and Stankovic, 196, 265

 Langhans, 275
 Lawson and Stewart, 342
 Leber, 33, 276, 248, 306
 Leclainche, 36, 433
 Leclainche and Vallée, 297, 433
 Leclef, 311, 312
 Leishmann, 313, 315, 329, 482
 Lemaire, 382
 Lemoine, 267
 Lenhartz, 473
 LePlay, 391
 Lepp, 74
 Lesné and Dreyfus, 375, 428
 Levaditi, 54, 200, 322, 426, 488
 Levaditi and Inmann, 316, 323
 Levaditi and Yamanouchi, 204
 Levin, I., 100
 Lewin, 461
 Lewis, 364, 374, 380, 382, 390
 Lidforss, 286
 von Liebermann, 175
 Liefmann, 173, 174, 183
 Liefmann and Cohn, 145, 176, 182
 Liesenberg and Zopf, 20
 von Lingselsheim, 136, 178, 472
 Linossier and Lemoine, 267
 Lister, 79, 134
 Loeb, 292, 517
 Loeb, Strickler and Tuttle, 406, 407
 Löhlein, 314
 Longcope, 303, 304
 Löw, 287
 Löwenstein, 441, 443
 Löwi and Meyer, 408, 409
 Lubarsch, 80, 81
 Lüdke, 477
 Lura, 402
 Lustig, 480
 Lustig and Galleotti, 70

 Macdonald, 319
 Macfadyn, 71, 222, 477
 MacKonky, 480
 Madsen, 39, 107, 113, 116, 117, 120, 121, 125, 130, 337, 338, 408
 Magendie, 359, 405
 Magnus, 255
 Malory and Wright, 353
 Malvoz, 224, 225
 Manwaring, 167, 205, 370, 404
 Maragliano, 148
 Marbé, 173
 Marchand, 297, 312, 325
 Marfan and LePlay, 391
 Marie and Levaditi, 200
 Marie and Morax, 41
 Marie and Tiffeneau, 133
 Marinesco, 41
 Markl, 479
 Markl and Rowland, 469
 Marks, 183, 460
 Marmorek, 469, 472, 473
 Marshall, 157
 Martin, 190, 193, 316, 321, 453
 Martin and Cherry, 105, 106
 Martini, 479
 Marx, 69, 100, 301
 Massart and Bordet, 288

- Mathes, 477
 Matschinsky, 276
 Matuso, 149
 Mayer, 70
 McClintock and King, 375
 McIntosh and Fildes, 200, 210
 McKenzie, 196
 McNeill, Archibald, 208, 216
 Meakin and Wheeler, 342
 Meakins, 341
 Meier, 201, 204
 Meltzer, 434
 Mendel, 96
 Mennes, 312, 325
 Menzer, 473
 Mesnil, 170
 Metchnikoff, 31, 46, 78-81, 84, 87, 89-91, 130, 131, 136, 138, 140, 169, 170, 172, 218, 272-275, 276, 277, 296-305, 308, 484
 Metchnikoff and Besredka, 67, 68, 349
 Meyer, 42, 256, 320, 357, 408, 409, 440, 447, 477
 Meyer and Gottlieb, 30, 39, 40, 44, 45, 99, 409
 Meyer and Michaelis, 473
 Meyer and Overton, 44
 Meyer and Ransom, 41, 131, 447
 Michaelis, 180, 246, 251, 293, 473
 Michaelis and Rona, 495
 Michaelis and Schick, 462
 Michaelis and Skwirsky, 179, 181
 Miller, 92
 Mironoff, 472
 Mita, 366, 367, 415, 430, 432
 Morax, 41
 Moreschi, 69, 189, 190
 Morgenroth, 86, 87, 106, 132, 142, 143, 146, 147, 150-154, 157, 165, 306, 359, 465
 Morgenroth and Ehrlich, 173, 177
 Morgenroth and Sachs, 163, 165, 324
 Moro, 356, 391, 438, 441
 Moss, 148
 Mosser, 473
 Mouton, 274
 Moxter, 305
 Much, 285, 342
 Muir, 145, 190, 195
 Muir and Browning, 177
 Muir and Martin, 190, 192, 316, 321
 Müller, 44, 47, 55, 87, 173, 200, 228, 250, 257, 261, 262, 267, 268, 307, 324
 Müller, Friedrich, 307, 494
 Müller and Jochmann, 306
 Myers, 251
 Nathan, 418
 Naunyn, 405
 Neisser, 155, 156, 198, 199
 Neisser and Döring, 196
 Neisser and Friedemann, 242, 243, 244, 265
 Neisser and Sachs, 189, 191, 212, 213
 Neisser and Wechsberg, 92, 160-163, 165, 186, 191, 245
 Nencki, 39, 77, 83
 Nernst, 122
 Neufeld, 17, 71, 75, 80, 90, 169, 290, 306, 317, 333, 344
 Neufeld and Bickel, 323
 Neufeld and Cole, 468
 Neufeld and Dold, 413, 415, 417, 418, 421, 423
 Neufeld and Haendel, 183, 194, 221, 351, 474, 475
 Neufeld and Hüne, 316, 317, 321
 Neufeld and Rimpau, 76, 315, 320, 321
 Neufeld and Töpfer, 321, 322
 Neumann, 343
 Nichols, 203, 234
 Nicolle, 193, 224, 250, 380, 394
 Nicolle and Abt, 387, 394
 Nikati and Rietsch, 54
 Nissen, 80
 Noguchi, 47, 164, 174, 175, 181, 183, 195, 196, 200, 203, 208, 210, 306, 438, 465
 Nolf, 173, 178, 395
 Norris, 251, 252
 Northrup, 446
 Novy, 29, 30, 463
 Nuttall, 51, 79-81, 84, 87, 134, 137, 254, 255, 328
 Obermeyer and Pick, 249-251, 258, 261
 Odaira, 402
 Oden, Sven, 510
 Ohlmacher, 432
 Olmstead, 216, 217
 Ople, 306, 308, 341
 Oppenheim, 18
 Oppenheimer, 29, 128, 250, 493
 Orthenberger, 178
 Osborne, Mendel and Harris, 96
 Oschida, 490
 Ostenberg, 261
 Ostwald, 293, 502
 Ottenberg, 208, 238, 239
 Ottenberg and Epstein, 58
 Ottenberg and Kaliski, 149
 Ottenberg, Kaliski and Friedemann, 148
 Ottenberg and Thalheimer, 148
 Otto, 361, 374, 376, 380-382, 386, 387, 487
 Overton, 44, 47
 Paltauf, 90, 224, 226, 232
 Panum, 272
 Park, 230, 349, 462
 Park and Biggs, 324, 449-451
 Park and Krumwiede, 52
 Park and Throne, 457
 Park and Williams, 452, 453, 455
 Pasteur, 1, 16, 56, 63, 64, 65, 83, 128, 136, 345, 481, 489, 490
 Pasteur and Thuillier, 16
 Paul, 54
 Paul, Kraus and Levaditi, 488
 Pauli, 241, 246
 Pearce, 93, 291, 350
 Pearce and Eelsenbrey, 368, 369, 398
 Pearce, Karsner and Eelsenbrey, 373
 Pearson, 344
 Perrin, 504, 505
 Peterson, 424
 Petterson, 297, 303, 305, 308, 328
 Pfaundler, 223
 Pfeffer, 135, 285, 286
 Pfeiffer, 33, 38, 69, 87-89, 137-140, 191, 232, 289, 324, 366, 367, 373, 378, 392, 412, 421, 444, 445, 487
 Pfeiffer and Beck, 53
 Pfeiffer and Bessau, 476
 Pfeiffer and Finsterer, 445
 Pfeiffer and Friedberger, 190, 191
 Pfeiffer and Isaacs, 88, 137
 Pfeiffer and Kolle, 482
 Pfeiffer and Marx, 69, 100, 301
 Pfeiffer and Mita, 366
 Pfeiffer and Wassermann, 85, 87, 88
 Philipp, 118
 Physalix, 464
 Physalix and Bertrand, 75, 86, 105
 Physalix and Contejean, 100
 Pick, 29, 36, 70, 224, 249, 250, 251, 258, 261, 264
 Pick and Schwartz, 250, 371
 Pick and Yamanouchi, 371, 388
 von Pirquet, 264, 265, 266, 356, 390, 397, 440-444
 von Pirquet and Moro, 438
 von Pirquet and Schick, 27, 361, 426, 427, 428
 Plant, 199
 Plotz, 55
 Pollack, 41, 131
 Pollender, 1
 Ponfick, 405
 Porges, 19, 200, 236, 243, 265, 266
 Porges and Meier, 200, 201, 204
 Portier, 360
 Portier and Richet, 360
 Potter, 341, 344
 Potter, Ditman and Bradley, 341
 Pötl, 200
 Powell, 353
 Preiz, 19
 Pribram, 87
 Proeschner, 220
 Prudden, 80
 Prudden and Hodenpyl, 69
 Quincke, 294
 Rabe, 97
 Rabinowitch, 205
 Rankin, 427
 Ransom, 39, 41, 447
 Ranzi, 214, 367, 373, 445
 Reagh, 225, 231

- Rees, 353
Rehns, 225
Reid, 314, 341
Reim, 246
Rhumbler, 294
Ribbert, 234, 281
Richet, 37, 360, 380, 382, 387, 428
Richet and Hérincourt, 74, 360
Richet and Portier, 360
Richter, 237
Rietsch, 54
Rimpau, 76, 315, 320, 321
Ritz, 185
Ritz and Sachs, 182, 415
Rodet and Lagrifoul, 477
Roepke, 351, 356, 357
Roessle, 92
Roger, 89, 218, 472
Rolleston, 426, 427
Römer, 65, 101, 263, 441, 461, 462
Römer, Field and Teague, 518
Römer and Sames, 461
Römer and Somogyi, 461
Rommel and Herman, 478
Rona, 495
Rondoni, 201
Rosenau, 70, 108, 110, 351, 436, 454, 456, 457, 489
Rosenau and Amoss, 436
Rosenau and Anderson, 362, 365, 368, 372-382, 389, 390, 401, 410, 411, 426, 428, 430, 437, 463, 464
Rosenblatt, 442
Rosenow, 22, 319, 325, 326, 424, 472
Rosenthal, 148
Roser, 272
De Rossi, 225
Rouget, 289, 297
Roux, 66, 73, 125
Roux and Behring, 107
Roux and Vaillard, 104, 125, 130
Roux and Yersin, 32, 36, 72, 77
Rowland, 71, 469, 480, 487
Ruediger, 178, 314, 318, 395
Ruffer, 234
Ruppel, 356
Rusk, 268, 269
Russ, 381, 382, 388, 389, 391, 394, 396, 400
Russell, 319, 482-484
- Sachs, 87, 130, 153, 155, 163, 165, 166, 174, 176, 182, 189, 190, 201, 212, 213, 324, 415
Sachs and Altmann, 184
Sachs and Kyes, 175
Sachs and Rondoni, 201
Sachs and Teruuchi, 179
Sacquépée, 228
Salecker, 423
Salmon and Smith, 20, 72
Salomonsen and Madsen, 125, 130, 337, 408
Sames, 461
Samuely, 29, 30
Sanarelli, 85, 87
Sanpietro and Tesa, 214
Sauerbeck, 135, 136, 326
Sawtschenko, 18, 320
- Schattenfroh, 34, 36, 73, 87, 304, 305
Schattenfroh and Grasseberger, 458
Scheller, 227
Schereschewsky, 203
Schick, 27, 361, 426-428, 447-449, 462
Schidorsky and Reim, 246
Schittenhelm, 370
Schittenhelm and Weichardt, 403, 435
Schmidt, 204, 259-262
Schmiedeberg, 30
Schneider, 171, 303, 305
Schreiber, 459
Schucht, 199
Schultz, 365, 397, 398
Schultze, 241
Schürmann, 39
Schütze, 94, 185, 253, 255
Schwann, 1
Schwartz, 250, 371
De Schweinitz, 73, 87
Sears, 246
Sears and Jameson, 242
Selmi, 28, 77
Sewall, 464
Shattock, 237
Shepilewsky, 131
Shibayama, 19
Shiga, 219, 220
Siedentopf and Zsigmondy, 503
Stegert, 447
Simon, 332
Simm and Lamar, 333
Simm, Lamar and Bispham, 319, 332, 333
Simon and Thomas, 214
Skwirsky, 179, 181
Slatineau, 92
Sleeswijk, 367, 394
Smith, Alexander, 119
Smith, Henderson, 172, 449
Smith, Theobald, 9, 20, 350, 453, 454
Smith and Reagh, 225, 231
Smoluchowski, 505
Sobernheim, 64
Sohma, 263, 372, 436
Southard, 364, 371, 380, 382, 386-390, 394
Sharr, 465
Stahl, 286
Stankovic, 196, 265
Steinhardt, 368, 374, 377, 379
Stenitzer, 469, 477
Stern, 80, 209
Stewart, 342
Stillmarck, 106
Stimson, 490
Strauss, 25
Strauss and Gumalela, 52
Streng, 167
Strickler, 406, 407
Strong, 67, 486, 487
Sturili, 237
Surmont, 92
Svedberg, 505
Sweet, 45
Swift, 201
Syme, 96
Szymanowski, 402, 418
- Takaki, 46, 47, 130, 132, 133
Tamancheff, 486
Tarassewicz, 169, 301
Tarozi, 5
- Tavel, 473
Teague and Buxton, 518
Terry, 284, 461
Teruuchi, 179
Tesa, 214
Thalheimer, 148
Thomas, 57, 214
Thomsen, 444
Throne, 457
Thucydides, 61
Thullier, 16
Tiffeneau, 133
Tizzoni, 84
Todd and White, 148
Töpfer, 321, 322
Toussaint, 64, 74
Trapetznikoff, 299
Traube, 79, 134, 185, 496
Tschernorutski, 284, 343
Tschistovitch, 94, 248
Tenda, 319
Tsuruski, 180
Tunnicliff, 325, 341
Turro and Gonzales, 430
Tuttle, 406, 407
- Uhlenhuth, 70, 94, 196, 254, 255-257, 263, 267, 444
Uhlenhuth and Haendel, 263, 373, 395, 406, 407
Uhlenhuth and Weidanz, 254, 268
- Vaillard, 84, 104, 125, 130
Vaillard and Rouget, 289
Vaillard and Vincent, 289, 463
Vaillard, Vincent and Rouget, 297
Vallee, 36, 433
Van Bemmelen, 501
Van der Velde, 73, 87, 168, 473
Van Ingen, Philip, 432
Vaughan, 31, 38, 366, 385, 393, 402, 412, 413, 419, 424, 476
Vaughan, Cumming and Wright, 367
Vaughan and Novy, 29, 30
Vaughan and Wheeler, 393, 403, 419
Vedder, 177
DiVestea and Zagari, 42
Vincent, 289, 297, 463
Volk, 200, 223, 226, 233, 235, 236, 389
- Wadsworth, 17, 26
de Waele, 37, 48
Wagner, 81
Waldeyer, 272
Walker, 8, 18, 172, 228, 229, 303
Walker and Swift, 201
Washburn, 89, 218
Wassermann, 34, 73, 85, 87, 88, 100, 105, 106, 131, 133, 145, 155, 200, 210, 322, 349, 390, 408, 422, 423, 424, 469, 470
Wassermann and Bruck, 192, 198, 439, 440
Wassermann and Citron, 21, 101, 469
Wassermann, Meisser and Bruck, 198

- Wassermann, Neisser, Bruck and Schucht, 199
 Wassermann and Plaut, 199
 Wassermann and Schütze, 94, 253
 Wassermann and Takaki, 46, 130, 132, 133
 Webb, Williams and Barber, 15, 67
 Wechsberg, 92, 104, 155, 160-165, 186, 191, 245
 Wechseltmann, 210
 Weichhardt, 392, 403, 429, 435, 436
 Weichhardt and Schittenhelm, 370
 Weidanz, 254, 268
 Weigert, 129, 291
 Weil, 76, 205, 398
 Weil and Braun, 200
 Weill-Hallé and Lemaire, 382
 von Weimarn, 500, 501, 502, 509
 Welch, 17
 Wells, Gideon, 29, 293, 294, 389, 390, 393, 403
 Wendelstadt, 155
 Werbitzky, 404
 Werigo, 289
 Wernicke, 66, 73, 83, 84, 86, 104
 Western, 318
 Weygant, 200
 Wheeler, 393, 403, 419
 White, 148
 Whitfield, 341
 Widai, 90, 220
 Wiener, 88
 Wilde, 156, 195
 Williams, 15, 67, 452, 453, 455
 Windsor, 313, 328
 Wirth, 462
 Wladimiroff, 53, 222, 252, 463
 Wolf-Eisner, 38, 393, 412, 434, 440, 441
 Wood, Francis Carter, 210, 221
 Wright, 68, 80, 90, 313, 314, 315, 323-346, 350-353, 367, 482
 Wright and Douglas, 313, 314, 315, 316, 334-337
 Wright and Reid, 314, 341
 Wright and Windsor, 313, 328
 Yamanouchi, 204, 371, 373, 388, 442, 444
 Yersin, 32, 36, 72, 77, 478
 Yersin, Calmette and Borrel, 478
 Yersin and Roux, 479
 Young, 242, 266, 269, 287, 429, 499, *et seq.*
 Zagari 42
 Zeissler, 185
 Zimmermann, 202
 Zinsser, 7, 193, 305, 400, 407, 415
 Zinsser and Carey, 183, 278, 279, 307
 Zinsser and Dwyer, 22, 228, 402, 476
 Zinsser and Johnston, 196
 Zinsser and Ottenberg, 261
 Zinsser and Young, 269, 429
 Zopf, 20
 Zsigmondy, 503, 512
 Zupnik, 251

INDEX OF SUBJECTS

- Abderhalden, protective ferments of, 493.
See also under Ferments, protective, in animal body.
- Abscesses, secondary, caused by bacteria, 25
- Absorption theory in toxin-antitoxin reaction, 123
- Acne, opsonic index in vaccine treatment of, 339
- Adrenal cytotoxin, 92
- Agglutination, absorption experiments of Castellani on, 232
- acid, 246
value of, for differential purposes, 246-247
- action of salts in, 240
- agglutinability of bacteria in, in agglutinoids, 229
normal differences in, between strains of same species, 228, 229
- agglutinins in, 223, 224
absorption of, 232
complete, impossibility of, 232
heating of, 226
explanation of, 236
major, 229
normal, 233
explanation of, 234
qualitatively identical with "immune" agglutinins, 234
para or minor, 229
- agglutininogen in, 223, 224
effects of heating of, 226
localization of, in ectoplasmic layers of cells, 224
- agglutinoids in, 235
- biological relationship and, 230
- Bordet's explanation of, 240
- by means of cell body proper, 225
- by means of ectoplasmic substances of bacteria, 224, 225
- by means of flagella, 224, 225
- cataphoresis of bacteria in, 242, 243
- description of, 218
- effect of gelatin addition in mastic solution on, 244
- Ehrlich's interpretation of process of, 234
- Agglutination, Ehrlich's interpretation of, diagrammatic representation of, 235
- Eisenberg and Volk's interpretation of, 235
- Ficker's reaction in, 223
- flocculation of colloids and, 241
mechanism of, 241, 242
mutual, 242
group, 229
cause of, 230
diagnostic value of, 231
- hemagglutination analogous to, 236, 237
- history of, 218
- importance of electrolytes in, 240
- in colloidal solutions, inhibition zones in, 245
- in excess of agglutinin, colloid phenomenon and, 518
- in immune serum, 141
- in motile and non-motile bacteria, 224, 225
- in salt-free environment, by addition of organic substances, 245
- influence of immunization with different animal species on, 232
with different species of bacteria on, 232
- influence of salts on sensitized and unsensitized bacteria in, 243, 244
- experiments of Neisser and Friedemann in, 244
- inhibition zones in, 162, 236
- iso-agglutinins in, 237
grouping of, 237, 238
value of presence of, 239
- methods of, 218 *et seq.*
Bordet's, 223
Ficker's reaction in, 223
Gruber-Widal, 220
macroscopic, 219
microscopic, 220
Proescher's, 220
thread reaction of Pfaunder, 222, 223
- nature of, 223
- not associated with life of bacteria, 222, 223
- Agglutination of "agglutinin" bacteria, 243
of bacteria in active immunization, 89
of capsulated bacteria, 243
phenomenon of, 218
power of, alterations in, by cultivation in immune serum, 228
effect of heating on, 226
spontaneous alteration in, 227
pro-agglutinoïd phenomenon in, explained as protective colloid action, 236
pro-agglutinoïd zone in, 162
pro-agglutinoids in, 235
relation of flagellar mechanism to, 222
specificity of, 219, 229
diagnostic value of, in group reaction, 231
limitations to, 229
thread reaction of Pfaunder in, 222, 223
"two phase" theory of, 241
- Agglutination reaction, diagnostic use of, 220, 221, 222
- flagellar mechanism in, 222
- in diagnosis of dysentery, 221
of glands in horses, 222
of paratyphoid fever, 221
of pneumonia, 221
of streptococcus infections, 222
of typhoid fever, 220, 221
- nature of, 239 *et seq.*
precipitin reaction analogous to, 263
with dead bacteria, 223
- Agglutinins, 223
absorption of, in agglutination, 232
complete, impossibility of, 233
- bacteriotropins and, 321
- definition of, 89
- group:
major, 229
para or minor, 229
heating of, effects of, 226
explanation of, 236
"immune," 234
in hemolytic serum, 93
nature of, 224
normal, 91, 233

- Agglutinins, normal, explanation of, 234
 qualitatively identical with "immune" agglutinins, 234
 production of, 129
 Agglutinogen, 223
 effects of heating on, 226
 localization of, in cell body proper, 225
 in ectoplasmic layer of cells, 224
 in flagellar substance, 224, 225
 nature of, 224
 Agglutinoids, 235
 Aggressins, 326
 action of, 21
 obtaining of, by bacteria
 secretion of, by virulence, 20-22
 Albuminolytins, 193, 211
 Alexin, 137
 a combination of soaps and proteins, 175
 absence of, from aqueous humor of the eye, 170
 analogy between ferments and enzymes and, 176
 bactericidal powers of, 137
 definition of, 80
 dependence of, on concentration, 176
 extraction of, from leukocytes and lymphatic organs, 304 *et seq.*
 filtration of, 177
 in hemolysis, 144
 inactivation of, by salt, 178
 by salt-removal, 178
 by shaking, 185
 reversibility of, 184
 Gramentski's experiments on, 184, 185
 increase of, on clot, 172
 influence of salts on action of, 177, 178
 leukocytic origin of, 168, 169 *et seq.*
 multiplicity of, 154 *et seq.*
 Bordet's views on, 156
 Ehrlich's views on, 155
 nature of, 137, 154
 chemical, 174, 175
 physical, 177
 presence of, in blood plasma, 170-172
 in blood stream, 170-172
 in normal blood, Gengou's view of, 303, 304
 Metchnikoff's view of, 302
 other experimenters on, 303
 production of, in liver, 173
 in thyroid gland, 172
 varieties of:
 macrocytase, 169
 microcytase, 169
- Alexin fixation, 186
 albuminolytin in, 193
 identity of, with precipitins, 193
 writer's opinion on, 193, 194
 Bordet and Gengou's experiment on, 186, 187
 Bordet-Gengou phenomenon in, 188 *et seq.*
 Gay's experiments supporting, 190
 in diagnosis of infectious diseases, 188
 in diagnosis of syphilis, 188
 Moreschi's experiments supporting, 189
 Bordetscher Antikörper in, 194, 195
 by immune animal sera with their specific antigens, 189
 by protein and antiprotein sensitizers, 189
 distinguished from complement deviation, 186
 during hemolysis, nature of, 176
 Ehrlich's (schematic) conception of, 187
 experiments of, on syphilitic monkeys, 198, 199
 forensic tests in, 211
 in anaphylaxis, 394
 in determination of nature of unknown protein, 211
 delicacy of, 212, 213
 technique of, 212
 in diagnosis of glanders, 216
 in diagnosis of gonorrheal infections, 216
 in diagnosis of malignant neoplasms, 213
 von Dungern's method of, 214
 antigen production for, 214
 results of, 215
 technique of, 214 *et seq.*
 in diagnosis of syphilis in human beings, 199
 in Wassermann reaction, 198
 nature of, 192 *et seq.*
 non-specific, 195
 by heated normal serum, 196
 by lipoidal substances in tissues, 195
 by preserved normal serum, 196, 197
 by protein emulsion and other extracts, 196
 by unsensitized bacteria, 195
 of specific precipitates, 190 *et seq.*
 albuminolytin identical with, 193
- Alexin fixation of specific precipitates, albuminolytin identical with, writer's opinion on, 193, 194
 Gay on, 190
 Pfeiffer and Friedberger on, 190
 Sachs on, 191
 writer on, 193, 194
 practical applications of, 198
 precipitin reaction and, 190, 192
 Dean on, 194
 Gengou on, 192
 Neufeld and Haendel on, 194
 writer on, 193, 194
 specific antiprotein sensitizers in, 192
 with syphilitic serum in antigens from normal organs, 200
 Alexin splitting, 178
 Brand on, 179
 by method of Ferrata, 178, 179
 by method of Liefmann, 184
 by method of Sachs and Altmann, 184
 effect of acid reaction on fractions of, 181
 end-piece in, 179 *et seq.*
 mid-piece in, 179 *et seq.*
 heat sensitiveness of fractions of, 180
 interchange of fractions of, in different animals, 182
 is it the inactivation of the mid-piece? 183
 mid-piece only bound in Wassermann reaction, 181
 physical occurrence of fractions of, in blood, 181
 presensitized cell in, 180
 properties of fractions of, 179
 quantitative relations between fractions of, 182, 183
 Alexocytes, 168
 Alkali-albuminate precipitin, 260
 "Alkalinity theory" of immunity, 83, 84
 Amanita phalloides, specific antitoxin from, 96
 Amboceptor, 149
 Bordet's definition of, 159
 complementophile groups or polyceptors of (Ehrlich), 149, 156
 cytophile group of, 149, 152, 153
 multiplicity of, 150, 151, 154
 Ehrlich and Morgenroth on, 150, 151
 quantitative determination of, in immune serum, 160, 161
 specificity of, 150

- Amboceptor and complement, Ehrlich and Sachs's views on union of, 164, 165
 Noguchi's measurement of quantitative relations of, 164
 quantitative ratio between, 163, 164
 Amebæ, artificial, 294
 Anaphylactic antibody, relation of, to other antibodies, 400
 Anaphylactic intoxication, peptone poisoning and, 404
 Anaphylactic shock, 363 *et seq.* *See also* Anaphylaxis, clinical manifestation of
 effect of atropin and other drugs on, 379
 Anaphylactin, 386
 Anaphylatoxin, 22, 396, 413
 action of alexin in, 422
 with normal or inactivated immune serum, 422, 423
 with salt solution, 424
 inhibition of, by too vigorous and prolonged reaction, 417
 source of, 424
 Anaphylaxis, 358
 alexin fixation in, 394
 analogy of immediate reaction in serum sickness to, 427
 analogy of serum sickness to, 428
 analogy of tuberculin reaction to, 442
 anaphylactic poisoning, nature of, 403 *et seq.*
 from precipitates, 396
 proteid split products of Vaughan and Wheeler in, 403
 symptoms of, similar to peptone poisoning, 404
 Anderson and Schultz's work on, 365
 anti-anaphylactic state in. *See* Antianaphylaxis
 antigen in, intervals between administrations of, 376
 identity of sensitizing and toxic substances of, 389
 Doerr and Russ's work on, 389
 Wells's work on, 389
 nature of, 370
 path of introduction of, 373
 intracerebrally, 373
 intravenously, 374
 intra-intestinally, 374
 by feeding, 374
 by rectum, 375
 into large intestine, 375
 subcutaneously, 374
 quantity of, administered, 376 *et seq.*
- Anaphylaxis, antigen in, specificity of, 371
 degree of, 371
 organ, 372
 auto-sensitization in, 373
 species, 372
 two separate substances in, 388
 Doerr and Russ's experiments on, 388, 389
 Gay and Adler's experiments in, 388
 Pick and Yamanouchi's experiments in, 388
 Arthus' work on, 361
 asthma and, 434
 Auer and Lewis's work on, 364
 autosensitization in, 373, 437
 Besredka and Steinhart's work on, 374 *et seq.*
 Besredka's theory of, 387
 Besredka's work on, 375 *et seq.*
 Biedl and Kraus's work on, 365, 368, 369
 Bogomolez's work on, 371
 Calvary's work on, 369
 cell participation in, 390, 397 *et seq.*
 clinical manifestations of, 363
 in dogs, 368
 fall in blood pressure in, 368
 fall of temperature in, 369
 increase of lymph flow in, 369
 intestinal reaction in, 370
 lowered coagulability of blood in, 369
 in guinea pig, 363
 alexin reduction in, 367
 circulation symptoms in, 366
 effect of atropin, and other drugs on, 365
 fall of temperature in, 366
 fever in, 366, 367
 lowered coagulability of blood in, 367
 pulmonary emphysema in, 364
 respiratory symptoms in, 364, 365
 susceptibility of various breeds in, 368
 temporary diminution of polynuclear leukocytes in, 367
 in rabbits, 368
 clinical significance of, 426
 dependence of, on preceding inoculation, 360
 diagnostic uses of, 444
 diminution of alexin after anaphylactic shock in, 394
- Anaphylaxis, diminution of alexin after anaphylactic shock in, significance of, 395
 early work on, 359
 Flexner's work on, 359
 Friedberger's work on, 366
 Friedemann's experiments in, *in vitro*, 395
 fundamental principles of, 358
 Gay and Southard's arguments against antigen-antibody reaction theory of, 387
 Gay and Southard's theory of, 386
 Gay and Southard's work on, 364
 hay fever and, 434
 in serum therapy. *See* Serum sickness
 in sudden attacks of catarrhal nasopharyngitis and conjunctivitis, 435
 in vaccine therapy, 432
 incubation time in, 360, 376
 Lesné and Dreyfus' work on, 375, 376
 Magendie on, 359
 Manwaring's work on, 370
 mechanism of anti-anaphylaxis in, 401 *et seq.*
 desensitization in, 401
 specificity in, 403
 tolerance to anaphylactic poison in, 402
 Nicolle's theory of, 394
 organ specificity in, 436
 Otto's work on, 361
 Pearce and Eisenbreys' work on, 368, 369
 Pfeiffer's work on, 366
 phenomena related to, 405
 toxic action of normal sera, 405
 toxin hypersusceptibility, 407
 Pick and Yamanouchi's work on, 371
 quantitative methods applied to study of, 391
 Ranzi's work on, 367
 relation of alexin to, 394
 relation of antibodies of, to other antibodies, 400
 Richet and Héricourt's work on, 360
 Richet and Portier's work on, 360
 Rosenau and Anderson's work on, 362, 374 *et seq.*
 sessile receptors, theory of, 390
 specificity of, 362, 363
 sympathetic ophthalmia and, 437
 Theobald Smith phenomenon in, 361
 Theobald Smith's work on, 363
 toxic action of normal sera and, 405

- Anaphylaxis, toxin hyper-susceptibility and, 407
 transference of, 362, 379 *et seq.*
 true antigen-antibody reaction, 390
 tuberculin ophthalmoreaction and, 440
 tuberculin reaction and, 438. *See also* Tuberculin reaction
 tuberculin skin reaction and, 440
 Vaughan and Wheeler's theory of mechanism of, 393
 Vaughan and Wheeler's work on toxic fraction of protein molecule in, 393
 Vaughan's work on, 366, 367
 Weichhardt and Schittenhelm's work on, 369, 370
 with bacterial extracts, 363
 with normal serum, 362
 with proteins, 362
 Anaphylaxis, bacterial, 410
 anaphylatoxin formation in, 413. *See also* under Anaphylatoxin.
 difference of speed of reaction of sensitized and unsensitized bacteria in, 418
 endotoxin theory of production of, 412
 Friedberger's experiments in, 413, 414
 facts deduced from, 415
 effect of excess of bacteria administered in, 415
 effect of excess of sensitization on anaphylatoxin in, 415
 effect of too prolonged exposure in anaphylatoxin in, 416
 nature of bacterial infections and, 419 *et seq.*
 Neufeld and Dold's experiments in, 417
 serum anaphylaxis and, 411
 Vaughan's theory of bacterial splitting as cause of, 412, 413
 Anaphylaxis, passive, 379 *et seq.*
 Biedl and Kraus's work on, 383
 Doerr and Russ's work on, 380
 duration of, 381
 Friedemann's work on, 380, 381, 383
 Gay and Southard's work on, 380
 interval between injection of sensitized serum and injection of antigen in, 382
 Anaphylaxis, passive, methods of production of, 380, 381
 nature of reaction of, 382
 Nicolle's work on, 380
 Otto's work on, 380
 Richet's work on, 380
 Weill-Hallé and Lemaire's work on, 382, 383
 Anderson and Schultz's work on anaphylaxis, 365
 Anthrax, relative susceptibility of man and animals to, 53
 study of, in regard to resistance and immunity, 296
 vaccination against, history of, 64
 method of, 64
 Anthrax bacilli, attenuation of virulence of, 18
 virulence of, 15
 Anti-alexin, 157
 action of, 157
 Anti-amboceptor, 152, 153
 Anti-anaphylaxis, 362, 377
 Besredka's work on, 375 *et seq.*
 mechanism of, 401 *et seq.*
 desensitization in, 401
 tolerance to anaphylactic poison in, 402
 methods of producing, 377
 Besredka and Steinhart's methods, 377
 Rosenau and Anderson's methods, 378
 specificity of, 378, 403
 "Anti-antibodies," 147
 Antibodies, concentration of, in lymphatic organs in active immunity, 100
 in other organs in active immunization, 101
 in active immunization, 85
 locality of production of, dependent on locality of antigen concentration, 101
 normal, explanation of, 234
 origin of, 100
 specificity of, 85
 Antibody formation, body cell in, 125
 chemical nature of, 126
 chemical action of antigens in, 128
 in active immunity, removal of spleen and, 100
 mechanism of (side chain theory), 124
 by internal secretion of body cell, 125
 processes of metabolism and, 125
 overproduction of receptors in, 128
 Antibody formation, principles of, 94
 Anticomplement, 157
 action of, 157
 Anticytophile interpretation of anti-sensitization (Ehrlich and Morgenroth), 153
 controversy on, 153
 "Antiformin," 70
 Antigen-antibody reactions, 129. *See also* Toxin - antitoxin reaction.
 antibody production in body cells in, 130
 relationship between susceptibility of tissue and toxin-binding properties in, 131
 side chain theory in, 129
 specificity of, 97, 129
 variety of antibodies in, 129
 agglutinins, 129
 antitoxins, 129
 cytotoxins, 129
 precipitins, 129
 Antigenic properties of cells, relation of, to lipid constituents, 97
 Antigens, action of, 95
 active immunization with, analogies to drug tolerance, 99
 characteristics of, variations in, 98
 complex structure of, Ehrlich and Morgenroth's conception of, 151
 definition of, 35, 94
 "local" immunity in organs directly in contact with, 101
 locality of production of antibodies dependent on locality of concentration of, 101
 organ specificity of, 98
 protein nature of, 96
 specificity of, 97
 Anti-isolysins, 147
 "Antiricin," 85
 Antisensibilisin, 387
 Antisensitization, anticytophile interpretation of (Ehrlich and Morgenroth), 153
 controversy on, 153
 Antisensitizers, 152, 153
 non-specificity of, 154
 Antitoxic serum, direct effect of, on toxin, 104
 indirect protective action of, against toxin, 104
 "normal" serum of Behring, 107
 Antitoxin, chemical relations of, with toxin, 114
 definition of, 85, 86
 diphtheria. *See* Diphtheria antitoxin.

- Antitoxin, production of, 129
by true toxins, 35
snake venom, 464
effect of heat on, 105
specific, substances inciting, 86
standardization of, 463
by means of toxin, 107
guinea pigs used in, 108
tetanus, production of, 463
use of, in passive immunization, 86
Antitoxin unit, diphtheria, 107
Antivenin, 464
Arrhenius and Madsen on neutralization in toxin - antitoxin reaction, 120
Arthus, phenomenon of, 380
work of, on anaphylaxis, 361
Ascoli and Izar's work on meistagmin reaction, 496
Asiatic cholera, relative susceptibility of man and animals to, 53
Asthma, anaphylaxis and, 434
Atrepsie, 56
Attenuation of bacteria by chemicals, 66
by cultivation under pressure, 66
by drying, 65
by heating, 65
by passage through animals, 65
by prolonged cultivation above optimum temperature, 65
by prolonged growth on artificial media in presence of own metabolic products, 65
capsule formation in, 18
Auer and Lewis's work on anaphylaxis, 364
Autocytotoxins, 93
Autogenous vaccines, 351
"Autohemolysins," 146, 147
Auto-inoculation by massage in active immunization, 340
Autolysins, 146
Autosensitization in anaphylaxis, 373
Auxilysin, 167
Avian tuberculosis, relative susceptibility of animals to, 52
Bacillus botulinus, action of, 4
Bacteria, adaptation of, in body, 6, 7
agglutinability of, alterations in, 226
by cultivation in immune serum, 228
Bacteria, agglutinability of, caused by heating, 226
normal differences in, between strains of same species, 228, 229
spontaneous, 227
in agglutinoid, 229
agglutination in. *See under Agglutination.*
"agglutinin" bacteria, agglutination of, 243
aggressin secretion of, in body, and virulence of, 20-22
anti-opsonic properties of, and antichemotactic substances, 325
artificial cultivation of, 10
attenuation of, 18
methods of, 65. *See also under Attenuation.*
by laboratory manipulations, 17
capsulated, agglutination of, 243
virulence of, 326
capsule formation in, and virulence, 18
colloid phenomena and action in, 516
destruction of, by cytases in leukocytes, 301, 302
by exudates, 300
by phagocytes, 300
different strains of, variation in infection from, 15, 16
ectoplasmic hypertrophy of, in relation to virulence, 19, 20
effect of body temperature on invasive powers of, 12
effect of cultural adaptation of, on virulence, 12
effect of path of introduction of, on infection, 12-14
on virulence of, 12-14
effect of quantity of, introduced, on infection, 14
entrance of, into body tissues, 6
generalized action of, 24
growth of, within leukocytes, 298
in blood stream, 24
in localized infection, reaction to, 26
through accidental conditions, 25
in normal serum, resistance to phagocytosis of, 325
incubation of, 26
localized action of, 23
measurement of relative degrees of virulence of, 15
negative charge of, in suspension, 242
number of, introduced, and relative virulence, 15
Bacteria, occurrence of, 2
parasitic and saprophytic, classification of, 11
phagocytosis of. *See under Phagocytosis.*
relative virulence of different strains of same, 15, 16
resistance of, to phagocytosis, due to nonabsorption of opsonin, 326
resistance of living cell to, 6
secondary abscesses caused by, 25
selective action of, in localized infection, 25
selective lodgment of, in tissues, 40
sensitized, immunization with, 68
sensitized and unsensitized, influence of salts on agglutination of, 243, 244
similar conditions produced by different, 23
specificity of, and infection, 22
use of, in active immunization, 85, 87-89
variation in virulence of, when successively passed through animals, 16, 17
Bacterial anaphylaxis. *See Anaphylaxis, bacterial.*
Bacterial extracts, active immunization with, 69
extraction of bacteria for, by mechanical methods, 71
by permitting them to remain in fluid media, 70
Bacterial infections, conceived as reaction of body against a foreign antigen, 420
nature of (Friedberger), 419 *et seq.*
Bacterial precipitins, group reactions in, 251
diagnostic value of, 252
partial or minor, 252
Bacterial products, active immunization with, 72
Bactericidal properties of blood serum, 134
Bacterial proteins, 33
Bacterial toxins. *See also Toxins.*
action of, after distribution in body, 40
active immunization with, 72
chemical structure of, in relation to toxicity, 43
endotoxins. *See Endotoxins.*

- Bacterial toxins, lesions produced by, in course of excretion, 45**
 local injury by, 45
 nature of, 32
 nature of union of, to body cells, 44
 nerves attacked by, 41
 obtaining of, 32
 from dead cultures, 32
 from living cultures, 32
 physical relationship with body cells in action of, 44
 production of antitoxin by, 35
 selective action of, 40
 principles of, 43
 reasons for, 45
 selective localization of, 39
 specific distribution after introduction of, 40
 specific susceptibility of tissues to, 45
 chemicals inhibiting, 47
 true, 33
 analogy of, with enzymes, 36
 bacteria producing, 34
 characteristics of, 34
 heat sensitiveness of, 36
 incubation time of, 36
- Bactericidal powers of blood serum, 79**
 alexin in, 137
 nature of, 137
in vivo, 137
 cholera experiments in, 137
- Bactericidal substances, origin of, from leukocytes, 169 *et seq.***
- Bacteriolysins, agglutinins and, 321**
 in active immunization, 89
- Bacteriolysis, 137**
 extracellular theory of, 140
 heat a factor in, 138
 mechanism of, 138
 immunity conferred by, 137, 138
 in immune serum, 137, 138
 Bordet's findings in, 140, 141
 inactivation and reactivation in, 140
 Pfeiffer's phenomenon in, technique of, 138 *et seq.*
 specificity of protection of, 137, 138
 transference of power of, 137, 138
 leukocyte action in, 168
 leukocytes in, 140
- Bacteriotropins, bactericidal sensitizers and, 321 *et seq.***
 normal opsonins and, 320 *et seq.*
- Bacteriotropins, presence of, in immune sera without lysins, 322**
 specificity of, 321
 thermostability of, 320
- Bail's aggressin theory, 21, 67**
 classification of parasites, 11
- Bauer's modification of Wassermann test, 209**
- Baumgarten's osmotic theory of bactericidal powers of blood, 135**
 Kitasato and Behring, Wernicke, antibody theory in active immunity of, 84
- Besredka and Steinhardt's work on anaphylaxis, 374 *et seq.***
 on serum therapy of typhoid fever, 476
- Besredka's anti-endotoxic serum in treatment of typhoid fever, 476**
 method of administration of antitoxin, 431
 theory of anaphylaxis, 387
 vaccines in prophylactic immunization against plague, 487
 work on antianaphylaxis, 375 *et seq.*
- Biedl and Kraus's work on anaphylaxis, 365, 368, 369**
 on passive anaphylaxis, 383
- Blood, non-putrefaction of, 79, 134, 256**
 phagocytic activities of, in immunity, 79
- Blood plasma, cell-free, inhibition of bacterial growth in, in immunity, 79**
 presence of alexin in, 170-172
- Blood serum, agglutination in, immune, 141**
 alexin in, 137
 nature of, 137
 antibacterial powers of, in immunity, 79
 anti-isolysins in, 147
 autohemolysins in, 146, 147
 bactericidal and agglutinating powers of, Wright's studies of, 328
 bactericidal action of immune, 81
 alexin in, 137
 early theories regarding, 134
 in natural immunity, 79, 80
in vivo, 137
 cholera experiments in, 137
 mechanism of, 135
- Blood serum, bactericidal action of immune, mechanism of, assimilation theory of, 136**
 by chemically unfavorable environment, 135
 osmotic theory of, 135
- bacteriolysis in immune, 137**
 Bordet's findings in, 140, 141
 cholera experiments in, 137
 summary of facts in, 138
 heat a factor in, 138
 mechanism of, 138
 immunity conferred by, 137, 138
 inactivation and reactivation in, 140
 intracellular theory of, 138
 leukocytes in, 140
 Pfeiffer's phenomenon in, technique of, 138 *et seq.*
 specificity of protection of, 137, 138
 bacteriolytic powers of, transferable, 137
 cell-free, bacterial growth in, 81
 hemolysins in, 148
 hemolysis in immune, 141
 alexin or complement in, 144
 analogy of, to bacteriolysis, 142
 Bordet's work on, 141
 Ehrlich and Morgenroth on mechanism of, 142
 haptophore groups in, 142
 relation of antigen, amboceptor and complement in, 143-145
 work of Ehrlich and Morgenroth on, 143-144
 work of Liefmann and Cohn on, 145
 isohemolysis in, 146
 protective action of, against bacteria, 50
- Blood stream, presence of alexin in, 170-172**
- Body fluids, bactericidal powers of, in natural immunity, 80**
 Bogomolez's work on anaphylaxis, 371
- Borden's method of agglutination, 223**
- Bordet, explanation of, on agglutination, 240**
 findings of, on bacteriolytic power of immune serum, 140, 141
 on neutralization in toxin-antitoxin reaction, 122
 views of, concerning relation of antigen, amboceptor and complement, 158, 159

- Bordet, views of, concerning relation of antigen, ambocceptor and complement, action of complex in, 159
schematic representation, 159
- Bordet-Danysz phenomenon in neutralization in toxin-antitoxin reaction, 123
- Bordet and Gengou's experiment on alexin fixation, 186, 187
- Bordet-Gengou phenomenon in alexin fixation, 188 *et seq.*
- Gay's experiments supporting, 190
- Moresch's experiments supporting, 189
- Botulinus poisoning, action of, 41
- Botulinus toxin, 4
- Bouillon Filtre (Denys), 357
- Bovine colloid, 167
- Bovine tuberculosis, relative susceptibility of man and animals to, 52
- Brownian movement in colloids, 505
- Buchner on bactericidal power of blood in natural immunity, 80
- Calmette's investigations in snake poisons, 464 *et seq.*
- ophthlmo-reaction, 440
- Capsule formation in bacteria, by attenuation, 18
- virulence and, 18
- Calvary's work on anaphylaxis, 369
- Carriers, bacillus, 2
- Castellani, absorption experiments of, in agglutination, 232
- Catarrhal nasopharyngitis and conjunctivitis, sudden attacks of, anaphylactic nature of, 435
- Cell receptors, overproduction of, 152
- Cellular theory of immunity, 136
- Cerebrospinal meningitis, epidemic, serum therapy in, 469
- early investigations in, 469
- Flexner and Jobling's work on, 470
- Jochmann's investigations in, 469, 470
- Kolle and Wassermann's investigations in, 469
- nature of action in, 471
- results of, 470, 471
- standardization of serum in, 471
- Chantemesse's early experiments in serum therapy of typhoid fever, 475
- Chemotaxis, 285
- anaphylatoxin and, 291
- Engelmann's studies in, 287
- influence of bacteria in, 288
- influence of bacterial extracts in, 288, 289
- malic acid in, 286
- of slime-molds or myxomycetes, 286
- of spermatozoa of ferns, 286
- Pfeffer's technique in, 286
- physical explanations of, 292 *et seq.*
- selective, 291, 295
- surface tension in, 293
- "artificial amebæ" and, 294
- Chicken cholera, vaccination against, history of, 63
- Cholera, active prophylactic immunization against, 484
- Ferran's early investigations in, 484, 485
- Haffkine's method in, 485
- Kolle's method in, 486
- Strong's method in, 486
- Asiatic, relative susceptibility of man and animals to, 53
- chicken, vaccination against, history of, 63
- effect of path of introduction of bacteria of, on infection, 14
- experiments in, showing bactericidal powers of blood serum in, 137
- hog, immunization with bacterial products in, 72
- Cobra antitoxin, action of, 465
- standardization of, 466
- Cobra lecithid, 175
- Cobra venom, action of, 465
- Coctoprecipitin, 258
- "Coefficient of extinction," 332
- Cole's work on serum therapy of pneumonia, 475
- Colloids, 499
- application of phenomena of, in electrical field, 518
- to action in animal body, 516
- to action in bacteria, 516, 517
- to action with eggs of Fundulus, 517
- to biology, 515 *et seq.*
- to Danysz toxin-antitoxin phenomenon, 517
- to nonagglutination in excess of agglutinin, 518
- chemical properties of, 508, 509
- Colloids, chemical properties of, chemical composition in, 508
- chemical reactions in, 508
- electrochemical ionization in, 509
- classification of, 500
- definition of, 499
- emulsion, 500, 510
- flocculation of, by electrolytes, 509
- acids and alkalis in, 509
- concentration of electrolyte in, 509
- explanation of, 510
- nature of sol in, 510
- precipitin reaction analogous to, 265
- salts in, 509
- zone phenomenon in, 511
- gel, 506
- Graham's work on, 499
- irreversible, 500
- lyophobic, 515
- lyophilic, 515
- mutual reactions of, 511
- in two oppositely electrical sols, 512
- explanation of, 513
- in two similarly electrical sols, 511
- protective action of electrolyte in, 512
- protective action of great excess of one colloid over the other, 512
- explanation of, 512
- nature of, 500
- physical properties of, 501 *et seq.*
- Brownian movement of particles in, 505
- distribution of particles in, 505, 506
- electrical properties in, 506
- form of particles in, 502
- kinetic energy in, 505
- size of particles in, measurement of, 502 *et seq.*
- microscopic, 503
- osmotic pressure in, 503
- rate of settlement in, 504
- ultrafiltration method in, 504
- surface tension in, 506
- et seq.*
- preparation of solutions of, 514, 515
- reaction in, analogous to complement deviation phenomenon of Neisser-Wechsberg, 162
- inhibition zones in, 162
- reversible, 500
- sol, 500
- stability of, 501
- suspension, 500, 510
- Complement. *See also* Alexin.
- amboceptor and, Noguchi's measurement of quantitative relations of, 164

- Complement, amboceptor and, quantitative ratio between, 163, 164
 union of, Ehrlich and Sachs's views on, 164, 165
 definition of, 144
 in hemolysis, 144
 multiplicity of, 154 *et seq.*
 Bordet's views on, 156
 Ehrlich's views on, 155
 nature of, 154
 chemical, 174
 Complement deviation, 160 *et seq.*
 argument in favor of Bordet's views, 162, 163
 colloid reactions, analogous to, 162
 Gay's explanation of, 163
 in hemolytic reactions, 163
 Morgenroth and Sachs's experiments supporting, 163
 pro-agglutinin zone reaction analogous to, 162
 Complement fixation, 186.
See also Alexin fixation.
 in determination of nature of unknown protein, 211
 delicacy of, 212, 213
 technique of, 212
 practical applications of, 198
 test of, in diagnosis of glanders, 216
 in diagnosis of gonorrheal infections, 216
 in diagnosis of malignant neoplasms, 213
 von Dungern's method of, 214
 antigen production for, 214
 results of, 215
 technique of, 214 *et seq.*
 Complement splitting, 178.
See also under Alexin, splitting of.
 Complementoid, 158
 Complementophile group of amboceptor, 149
 Conglutinin, 167
 Conjunctiva, susceptibility of, to infection, 13
 Corpus luteum cytotoxin, 92
 "Cryptogenic tetanus," 5
 Cultivation of bacteria, artificial, 10
 Cytases, in phagocytes, destruction of bacteria by, 301, 302
 Cytolysins, 92
 Cytolytic substances, origin of, from leukocytes, 169 *et seq.*
- Cytophile group of amboceptor, 149, 152, 153
 Cytotoxins, 92
 specificity of, 92
- Danysz effect in neutralization in toxin-antitoxin reaction, 123
 Danysz toxin-antitoxin phenomenon, application of colloid phenomena to, 517
 Daphnia, Metchnikoff's study of, 274, 296
 phagocytosis in, 296
 Dean's antiplague sera, 480
 Denys and Leclef on importance of phagocytosis in immunity, 311, 312
 Diphtheria, active immunization in, with toxin - antitoxin, 458
 relative susceptibility of man and animals to, 53
 Diphtheria antitoxic serum, normal, 107
 Diphtheria antitoxin, 446
 antitoxin production in, 455
 concentration of, 457
 presence of, in blood of normal individuals, 448
 preservation of, 108
 speed in administration of, 447
 speed in absorption of, 449
 on intravenous injection, 449, 450
 on subcutaneous injection, 449, 450
 speed of diagnosis for, necessity of, 451
 stability of, 108
 standardization of, 455
 by means of toxin, 107
 early attempts in, 107
 statistics showing reduction of mortality with, 446
 toxin production for, 452
 choice of culture in, 452
 cultivation of strain in, 452, 453
 culture medium in, 453
 "maturing" of toxin in, 453
 testing of toxin in, 454
 Theobald Smith's method of, 454
 unit of, 107
 Diphtheria bacillus, action of, 4, 5
 Diphtheria toxin, action of, 40
 construction of, 118
 determination of diphtheria immunity with, 462
 normal, 107
 stability of, 108
- Diphtheria toxin, unit of, 107
 Diphtheria toxin-antitoxin, neutral mixtures of, 458
 Behring's method of immunization with, 458
 advantages of, 459
 chief value of, 459
 danger of anaphylaxis in, 459
 determination of presence of free toxin or antitoxin in convalescents following treatment with, 462
 human susceptibility to, 459
 production of homologous antitoxin in human beings with, for passive immunization, 460
 results of treatment with, 460
 standardization of antitoxin, 460, 461
 limes-necrosis of toxin in, 461
 Römer's method of, 461
 toxic action of, 458
 Doerr and Russ's experiments on two separate substances in anaphylactic antigen, 388
 Doerr and Russ's work on passive anaphylaxis, 380
 Dochez and Gillespie's work on serum therapy in pneumonia, 475
 Drug tolerance, analogy between, and active immunization with antigens, 99
 Dunbar's work on hay fever, 434
 von Dungern's method of alexin fixation in diagnosis of malignant tumors, 214
 antigen production in, 214
 results of, 215
 technique of, 214 *et seq.*
 "Dust cells," of the lungs in phagocytosis, 279
 Dysentery, agglutination reaction in diagnosis of, 221
- Ehrlich, conception of alexin fixation (schematic) of, 187
 of relation of antigen, amboceptor and complement, 149, 150
 interpretation of agglutination by, 234
 diagrammatic representation of, 235

- Ehrlich, on multiplicity of alexin or complement, 155
side chain theory of, in toxin - antitoxin reaction, 124
- Ehrlich's "antiricin," 85
- Ehrlich and Morgenroth on multiplicity of amboceptor, example of, 150, 151
- Ehrlich-Sachs phenomenon in sensitization, 165
- Bordet and Gay's interpretation of, 166, 167
- Eisenberg on residue antigen and antibody in precipitin reaction, 268
- Eisenberg and Volk's interpretation of agglutination, 235
- Endocarditis, malignant, 24
- Endolysins, 305
- Endotoxins, 33, 34
characteristics of, 37
toxic cleavage products of, 38
- Engelmann's studies in chemotaxis, 287
- Enzymes, analogy of, with true bacterial toxins, 36
in phagocytosis, endocellular and extracellular, 305
- Epithelioid cells, action of, in phagocytosis, 284
- Epitoxoids, definition of, 112
- Erythrocyte laking, 91
- "Exhaustion theory" of immunity, 83
- Exotoxins, 33. *See also* Toxins, true.
bacteria producing, 34
characteristics of, 34
chemically indefinable nature of, 35
- Fermentation, infectious disease and, 1
micro-organisms causing, 1
- Ferments, protective, in animal body, 493
- Abderhalden's experiments with, 494
significance of, 495
diagnostic value of, in pregnancy, 496
difference of, from antibodies, 495
leukocyte origin of, 494
methods of determining presence of, in blood, 494
dialysis method, 494
optical method, 494
- Ferran's investigations in active prophylactic immunization against cholera, 484, 485
- Ferrata, experiments of, in complement splitting, with salt solution, 179
- Ficker's reaction in agglutination, 223
- Flexner's observations on anaphylaxis, 359
on serum therapy in cerebrospinal meningitis, 470
- Forensic alexin fixation tests, 211
- Forensic determination of unknown proteins, 211
delicacy of, 213
technique of, 212
- Fornet and Müller, ring test of, for precipitin blood tests, 257
- Friedberger, experiments of, in bacterial anaphylaxis, 413, 414
on the nature of bacterial infections, 419 *et seq.*
work of, on anaphylaxis, 366
- Friedemann, experiments of, on anaphylaxis *in vitro*, 395
work of, on passive anaphylaxis, 380, 381
in rabbits, 383
- Fundulus, Loeb's experiments with eggs of, 517
- Garbat and Meyer's work on serum therapy of typhoid fever, 477
- Gastric juice, action of, on stomach itself, 6
- Gastro-toxin, 92
- Gay and Adler's experiments on two separate substances in anaphylactic antigen, 388
- Gay and Southard's objections to antigen-antibody theory of anaphylaxis, 387
theory of anaphylaxis, 386
work on anaphylaxis, 364
on passive anaphylaxis, 380
- Gay's sensitized killed vaccines in prophylactic typhoid fever immunization, 484
- Gels, 500
- Giant cells in phagocytosis, 280
foreign body, 280
tuberculous, 280
- Glanders, alexin fixation test in diagnosis of, 216
in horses, agglutination reaction in diagnosis of, 222
relative susceptibility of man and animals to, 53
- Gonococcus, relative susceptibility of man and animals to, 53
- Gonorrheal infections, alexin fixation test in diagnosis of, 216
- Gottstein-Mathes' work on serum therapy of typhoid fever, 477
- Graham's work on colloids, 499
- Gramenitski's experiment in reversal of alexin inactivation by heating, 184, 185
- Grobman on inhibition of bacterial growth by cell-free blood plasma in immunity, 79
- Gruber-Widal reaction in diagnosis, 220
- Haffkine's early work on prophylactic immunization against plague, 486
method in active prophylactic immunization against cholera, 485
- Haptines, 129
of the third order, 150
varieties of, 129
- Haptophore group in toxin, 110
action of, 110
- Haptophore groups in hemolysis, 142
- Hay fever, anaphylaxis and, 434
Dunbar's study of, 434
reaction in, 434
anaphylactic nature of, 435
toxic nature of, 435
treatment of, 435
- Heat-alkali-precipitin, 259
- Heat-precipitins, 259
- Hemagglutination, agglutination of bacteria by serum analogous to, 236, 237
- Hemoglobinuria, paroxysmal, 147
autohemolysis in, 147
hemolysis in, 147
- Hemolysinogens, human, 148
nature of, 148
- Hemolysins, anti-isolysins in, 147
autolysins in, 146, 147
isohemolysins in, 146
specific, definition of, 92
specific inciting of, in animal, 91
- Hemolysis, alexin or complement in, 144
amboceptor in, action of, 149 *et seq.*
anti-amboceptor in, 152, 153
anti-cytophile interpretation of anti-sensitization in (of Ehrlich and Morgenroth), 153
controversy on, 153
antisensitizer in, 152, 153
anti-isolysins in, 147

- Hemolysis, experiments of
Liefmann and
Cohn on, 145
in immune serum, 141
analogy of, to bacteri-
olysis, 142
Bordet's work on, 141
Ehrlich and Morgen-
roth on mechan-
ism of, 142
haptophore groups in,
142
relation of antigen,
amboceptor and
complement in,
143-145
multiplicity of ambocep-
tor in, 150, 151,
154
recapitulation of views
of Ehrlich and
Morgenroth on,
152
- Hemolytic properties of
normal serum, 91
- Hemolytic reactions, com-
plement deviation
in, 163
- Hemolytic serum, agglu-
tinins in, 93
Ehrlich and Morgen-
roth's conception
of neutralization
of, by antilysin
or anti-ambocep-
tor reacting with
cytophile group,
153
precipitins in, 94
- Hemolytic substances, ori-
gin of, from leu-
kocytes, 169 *et*
seq.
- Hepatotoxin, 92
- Hiss, investigations of, on
therapeutic use of
leukocyte ex-
tracts, 309, 310
- Hog cholera, immunization
with bacterial
products in, 72
- Högyes
method of treat-
ment in rabies,
492
- Holobut's work on bac-
terial anaphy-
laxis, 411
- Hopkins' method of stand-
ardization of vac-
cines, 353
- "Horror autotoxicus," 147
- Human isolysins. *See* Iso-
lysins, human
- Humoral theory of immu-
nity, 136
- Hydrophobia, active pro-
phylactic immu-
nization against,
489. *See also*
under Rabies
- Hypersusceptibility. *See*
Anaphylaxis
toxin, and anaphylaxis,
407
- Immune serum. *See also*
Serum, immune.
agglutination in, 141
bacteriolytic power of,
transferable, 137
bacteriolytic properties
of, Bordet's find-
ings in, 140, 141
- Immune serum, direct
neutralization of
toxin - antitoxin
reaction, the pro-
tective power of,
124
- hemolysis in, 141
alexin or complement
in, 144
analogy of, to bac-
teriolysis, 142
Bordet's work on, 141
Ehrlich and Morgen-
roth on mechan-
ism of, 142
haptophore groups in,
142
relation of antigen,
amboceptor and
complement in,
143-145
work of Ehrlich and
Morgenroth on,
143-144
work of Liefmann
and Cohn on,
145
- phagocytosis in, 90
specific agglutination of
bacteria in, 89
precipitin formation
in, 90
- Immune isolysins, 148
- Immunitäts Einheit, 107
- Immunity, acquired, 60
artificially, 63
definition of, 62
history of, 61
increased phagocytosis
and, 299
active, relation of pha-
gocytosis to, 329
cellular theory of, 136
definition of, 3
diphtheria, determina-
tion of, with
diphtheria toxin,
462
"high tide" of, 340
humoral theory of, 136
lasting, diseases in
which one attack
conveys, 60
diseases in which one
attack does not
convey, 61
local, in organs directly
in contact with
antigens, 101
in skin infections, 102
natural, 49
cellular theory of, 80
definition of, 50, 62
humoral theory of,
80
inflammation in, 78
mechanism of, 78
theories concerning,
78-82
bacterial destruc-
tion by phago-
cytic cells, 78
bacterial growth
in cell-free
blood serum, 81
bactericidal pow-
er of blood in
natural immu-
nity, 80
bactericidal pow-
er of normal
blood in nat-
ural immunity,
79
- Immunity, natural, mech-
anism of, theories
concerning, bac-
tericidal proper-
ties of extra-
vascular plasma
or serum, 81
inhibition of bac-
terial growth
by cell-free
blood plasma,
79
intracellular de-
struction of
bacteria, 78
phagocytic activi-
ties of blood,
79
phagocytic activities
of blood in, 79
principles of, 50
body temperature
in, 51
cultural conditions
for bacteria in
body a factor in,
56
increased invasive
powers of bac-
teria in, 56
individual differ-
ences in, 58
inheritance in, 56
racial differences in,
55
relative resistance
of animals in, 51
species resistance
in, 51
Pfeiffer phenomenon in,
138
phagocytosis in, 90
immunization, 60
against snake venoms,
464
history of, 61
immunization, active. *See*
*also Vaccine ther-
apy.*
against anthrax, 64
against chicken cholera,
63
against small-pox, 62
agglutination of bacteria
in, 89
"alkalinity theory" of,
83, 84
antibacterial, 85, 87
antibodies in, 85
bodies in, fundamen-
tal principles of
theory of, 84
origin of, 100
antitoxic, 85
as a therapeutic meas-
ure, action of, in
generalized sys-
temic infections,
347
in local infections,
346
in successive local
infections, 347
value of, 346
in acute diseases,
350
in subacute or
chronic cases, 349
as prophylactic measure,
value of, 345, 346
autogenous vaccines in,
351
auto-inoculation by mas-
sage in, 340

- Immunization, bacteria
 - used in, 85, 87-89
 - bacteriolysins in, 89
 - by means of living but attenuated cultures, 65
 - concentration of antibodies in lymphatic organs in, 100
 - in other organs in, 101
 - definition of, 63
 - "exhaustion theory" of, 83
 - "high tide" of immunity in, 340
 - in diphtheria, with toxin-antitoxin mixtures, 458. *See also* Diphtheria toxin-antitoxin.
 - invasion of bacteria in, mechanism of reaction in tissue cells against, 102
 - locality of production of antibodies dependent on locality of antigen concentration in, 101
 - negative phase in, 338
 - second injection in, 338
 - successive inoculations in, 338, 339
 - summation of, 338, 339
 - non-bacterial antitoxin-stimulating substances in, 86, 87
 - "osmotic theory" of, 84
 - phagocytosis in, 90
 - phenomena following, 82
 - precipitin formation in, 90
 - reaction of tissue cells to invasion in, 102
 - removal of spleen in, and antibody-formation, 100
 - "retention theory" of, 83
 - second positive phase in, 339
 - specificity of antibodies in, 85
 - summation of positive phase in, 339
 - tuberculin in, 355
 - vaccines in, 351
 - production of, 351
 - sensitized, 355
 - with dead bacteria, 351
 - with living bacteria, 351
 - standardization of, 353
 - Hopkins' method of, 353
 - Wright's method of, 352, 353
 - with antigens, analogy between drug tolerance and, 99
 - with bacterial extracts, 69
 - extraction of bacteria for, by mechanical methods, 71
- Immunization, with bacterial extracts,
 - extraction of bacteria for, by permitting them to remain in fluid media, 70
 - with bacterial products, 72
 - with dead bacteria, 68
 - methods used in killing bacteria for, 68
 - with fully virulent cultures in sublethal amounts, 66-68
 - with sensitized bacteria, 68
- Immunization, active prophylactic, in man, 481
 - against cholera, 484. *See also under* Cholera
 - against plague, 486. *See also under* Plague
 - against rabies, 489. *See also under* Rabies
 - against small-pox, 488. *See also under* Small-pox
 - against typhoid fever, 482. *See also under* Typhoid fever
- Immunization, passive, 74
 - antitoxins in, 86
 - definition of, 64
 - history of, 74
 - in diphtheria. *See* Diphtheria antitoxin
 - in diseases caused by bacteria which do not form soluble toxins, 466. *See also under* Serum therapy.
 - therapeutic application of, 75
 - toxin-antitoxin reaction in, 104
 - underlying principles of, 75
- Immunized animals, bacteriolysis in, 137
 - summary of facts in, 138
- Incubation of bacteria, 26
- Infection, acquired resistance to, 60
 - adaptation of bacteria in tissues in, 6, 7
 - aggressin secretion of bacteria in body and, 20-22
 - body temperature and, 51
 - capsule formation of bacteria and, 18
 - chronic, adaptation of bacteria in, 8
 - clinical manifestations of, 28
 - conjunctiva susceptible to, 13
 - criteria governing, 3
 - cultural conditions for bacteria in body and, 56
 - defence of intestinal tract in, 12
 - defence of mucous membranes in, 12, 13
 - defence of skin in, 12
- Infection, definition of, 5, 6
 - different, produced by same bacteria, 23
 - effect of body temperature on invasive powers of bacteria in, 12
 - effect of cultural adaptability of bacteria on, virulence of, 12
 - effect of path of introduction of bacteria on, 12-14
 - effect of quantity of bacteria introduced on, 14
 - entrance of bacteria in body tissues in, 6
 - focus in, 7-9
 - from bacteria in blood stream, 24
 - generalized, 24
 - increased invasive powers of bacteria a factor in, 56
 - incubation of bacteria in, 26
 - individual differences and, 56
 - inheritance and resistance to, 56
 - localized, 23
 - reaction in, 26
 - selective action of bacteria in, 25
 - through accidental conditions, 25
 - natural resistance against, 49
 - of various diseases, relative susceptibility of man and animals to, 52
 - protective action of blood serum against, 50
 - protective action of leucocytes against, 50
 - protective action of tissues against, 50
 - ptomains and, 28
 - ptomains as indirect cause of, 31
 - racial differences and, 55
 - resistance of living cell to, 6
 - secondary abscesses in, 23
 - secondary modifying factors in, 2
 - selective lodgment of bacteria in body and, 40
 - similar, produced by different bacteria, 23
 - species resistance to, 51
 - specificity of bacteria and, 22
 - susceptibility to, racial differences in, 55
 - relative, 51
 - variation in, of different strains of same bacteria, 15, 16
 - variation in degree of, in bacteria successively passed through animals, 16, 17

- Infection without infectious disease, 6, 7
 Infectious disease, definition of, 6, 8
 Inflammation, process of, and phagocytosis, 280 *et seq.*
 with pyogenic staphylococci, 281
 with tubercle bacilli, 283
 Influenza, relative susceptibility of man and animals to, 53
 Inheritance, a factor in resistance to infection, 56
 iso-agglutinins in blood serum influenced by, 58
 Inhibition zones in colloid reactions, 162
 in precipitation and agglutination, 162
 of sera in agglutination, 236
 Intestinal tract, defence of, in infection, 12
 Iso-agglutinin, 237
 grouping of, 237, 238
 in blood serum, 58
 value of presence of, 239
 Isohemolysins, 146
 Isolysins, human, 148
 grouping of, 148
 testing of, for transfusion, 149
 iso-agglutinins analogous to, 237
 Isoprecipitins, 255
- Jacobsthal's ultramicroscopic method of finding precipitates in syphilitic sera, 204
- Jenner, Edward, experimentation of, for immunization against small-pox, 62
- Jobling's work on serum therapy in cerebrospinal meningitis, 470
- Jochmann's investigations in serum therapy of epidemic cerebrospinal meningitis, 469, 470
- Kolle's method of prophylactic vaccination in cholera, 486
- Kolle and Otto's investigations in prophylactic immunization against plague, 487
- Kolle and Wassermann's investigations in serum therapy of epidemic cerebrospinal meningitis, 469
- Kraus, Rudolf, discovery of specific precipitins by, 248
- Kraus and Doerr's study of bacterial anaphylaxis, 410, 411
- Kraus and Stenitzer's serum in treatment of typhoid fever, 477
- L., definition of, 109
 method of determination of, 109
 L., definition of, 109
 constancy of, 110
 method of determination of, 110
- Laking, erythrocyte, 91
 "Landsteiner phenomenon" of autohemolysis in hemoglobinuria, 147
- Leishmann's technique for determination of opsonic index, 329
- "Leistungskern," definition of, 126
- Leprosy, relative susceptibility of man and animals to, 54
- Lcsné and Dreyfus' work on anaphylaxis, 375, 376
- Leukine, 305
- Leukocyte extracts, therapeutic use of, 308 *et seq.*
- Leukocytes, alexin extraction from, 304 *et seq.*
 growth of bacteria in, 298
 in bacteriolysis, 140
 action of, 168
 in leukocytosis, action of, 275, 276
 in phagocytosis, 324
 origin of bactericidal and hemolytic substances from, 168, 169 *et seq.*
 phagocytic powers of, 50
 proteolytic enzymes from, in phagocytosis, 306, 307
- Leucocytosis, 290
 bacteria decreasing, 290
 bacteria increasing, 290
 sources of leukocytes in, 290
- Leukoproteases, 306, 307
- Leukotoxin, 92
- Limes-necrosis (L-n), 461
- Lipoid constituents of cells, relation of, to antigenic properties, 97
- Lister on phagocytic activities of blood in natural immunity, 79
- Liver, production of alexin in, 173
- Loeb's experiments with eggs of *Fundulus*, 517
- Lubarsch on bactericidal properties of extravascular plasma or serum in immunity, 81
- Lüdke's work on serum therapy in typhoid fever, 477
- Lustig's antiplague serum, 480
- Lysins, production of, 130
- Macrocytase, 169, 301
- Magendie on anaphylaxis, 359
- Malta fever, relative susceptibility of man and animals to, 53
- Manwaring's work on anaphylaxis, 370
- Markl's serum in treatment of plague, 479
- Marmorek's work on serum therapy of streptococcus infections, 472, 473
- Measles, relative susceptibility of man and animals to, 54
- Meat poisoning, 4, 31
- Meistagmin reaction, 496
- Ascoli and Izar's experiments in, 496, 497
 value of, in diagnosis, 497
- Meningitis, epidemic cerebrospinal, serum therapy of, 469.
See also under Cerebrospinal meningitis, epidemic
- Metabolism, processes of, compared with those of antibody formation, 125
- Metchnikoff and Besredka's living sensitized vaccines for prophylactic typhoid immunization, 484
- Metchnikoff on bacterial growth in cell-free blood serum, 81
 theory of, on bacterial destruction by phagocytic cells in natural immunity, 78
- Metchnikoff's soured milk therapy, 31
- Microcytase, 169, 301
- Minimum lethal dose, definition of, 108, 109
 method of determination of, 109
- M L D, definition of, 108, 109
 method of determination of, 109
- Morgenroth's toxin-HCl modification in toxin-antitoxin reaction, 106
- Mucous membranes, defence of, in infection, 12, 13
- Mushroom, specific antitoxin from, 96
- Narcotics, reduction of phagocytosis by, 299
- Natural immunity. *See* Immunity, natural
- "Negative" phase in active immunization, 338
 second injection in, 338
 successive inoculations in, 338, 339
 "summation" of, 338, 339

- Neisser and Friedemann, experiments of, on influence of salts on sensitized bacteria in agglutination, 244
- Neisser and Wechsberg, phenomenon of, 160 *et seq.*
- analogous to colloid reactions, 162
- argument in favor of Bordet's views, 162
- Gay's explanation of, 163
- Morgenroth and Sachs' experiments supporting, 163
- pro-agglutinoid zone reaction analogous to, 162
- Neoplasms, malignant, alexin fixation in diagnosis of, 213
- von Dungern's method of, 214
- antigen production for, 214
- results of, 215
- technique of, 214 *et seq.*
- Nernst on views of Arrhenius and Madsen on neutralization in toxin-antitoxin reaction, 122
- Neufeld and Dold's experiments in bacterial anaphylaxis, 417
- Neufeld and Haendel's work on serum therapy of pneumonia, 474
- Neurotoxin, 92
- in snake venom, 465
- Nicolle's theory of anaphylaxis, 394
- work on passive anaphylaxis, 380
- Noguchi's modification of the Wassermann test, 208, 209
- schematic presentation of, 209
- "Normal" diphtheria antitoxic serum, 107
- "Normal" diphtheria toxin, 107
- "Normal" serum. *See under* Serum
- agglutinins in, 91
- hemolytic properties of, 91
- opsonins in, 91
- toxic action of, and anaphylaxis, 405
- Nuttall on bactericidal power of normal blood in natural immunity, 79
- Nuttall's experiments on determining zoological classifications by means of precipitin reaction, 254, 255
- Ophthalmia, sympathetic, Elschinig's explanation of, as anaphylactic reaction, 437
- Opium, reduction of phagocytosis by, 299
- Opsonic action, phagocytosis due to, 313
- Opsonic index, determination of, Leishmann's technique for, 329
- Simon, Lamar and Bispham's technique of, 332, 333
- Wright's technique for, 330 *et seq.*
- difficulties in, 332, 333
- value of, 333
- fluctuation of, in untreated patients under influence of exercise of diseased parts, 340
- in autoinoculations by massage, 340
- in sera of normal and infected individual, comparison of, 334
- in serum therapy, comparison between that in exudate of infected foci and blood serum, 340
- in staphylococcus infections, 334
- during vaccine treatment with dead staphylococcus cultures, 335
- in treatment of gonorrheal arthritis with autoinoculation by massage, 340
- in vaccine therapy, improvement and, 341
- of acne, 339
- of staphylococcus furunculosis, 335
- of syccosis, 336, 337
- of tuberculosis, 341-343
- value of, in controlling therapeutic vaccinations, 344
- in showing degree and conditions in which vaccination is successful, 344, 345
- vaccine therapy and, 328 *et seq.*
- value of, in therapeutics, 338
- Opsonic powers of normal serum, 314
- reduction of, by heat, 314
- Opsonins. *See also* Phagocytosis
- definition of, 313
- immune, bactericidal sensitizers and, 321 *et seq.*
- increase of, 315
- heated, increase of power of, by addition of fresh normal serum, 318
- reactivation of, by addition of alexin, 318
- normal and, 320 *et seq.*
- Opsonins, immune, resistance of, to heat, 315
- specificity of, 321
- thermostability of, 320
- normal, 91
- coöperation of heat-stable and heat-sensitive body in, 319
- instability of, 314, 315
- nature of, 316
- similarity of, to alexin or complement, 316, 317
- specificity of, 318
- production of, in thyroid gland, 173
- qualitative difference between normal and immune, 315, 316
- specific thermostable, in normal serum, 317
- Organ specificity of antigens, 98
- "Osmotic theory" of immunity, 84
- Otto's work on anaphylaxis, 361
- in passive anaphylaxis, 380
- Pancreas cytotoxin, 92
- Panum's theory of intracellular destruction of bacteria in natural immunity, 78
- Parasites, biological transition of saprophytes to, 5
- Bail's classification of, 11
- Parasitic bacteria, 4
- Paratyphoid fever, agglutination reaction in diagnosis of, 221
- Paroxysmal hemoglobinuria, 147
- hemolysis in, 147
- Partial absorption method of Ehrlich in measurement of toxin-antitoxin combination, 115
- Pasteur, "exhaustion theory of," 83
- experimentation of, on immunization against chicken cholera, 63
- work of, on immunization against anthrax, 64
- on prophylactic immunization in rabbits, 489
- Pathogenic bacteria, adaptation of, in tissues, 6, 7
- entrance of, in body tissues, 6
- saprophytic nature of certain, 4
- Pathogenic microorganisms, definition of, 3
- occurrence of, 2
- resistance of living cell to, 6

- Pearce and Eisenbrey's work on anaphylaxis, 368, 369
- Persensitized cells, 180
- Petterson's investigations on therapeutic use of leukocyte extracts, 308
- Pfaundler's thread reaction in agglutination, 222, 223
- Pfeiffer on causes of bacterial anaphylaxis, 412
- work of, on anaphylaxis, 366
- "Pfeiffer phenomenon" in active immunization, 89
- in bacteriolysis, technique of, 138 *et seq.*
- Metchnikoff's view of phagocytosis in peritoneal exudate and, 302
- Phagocytes 276
- fixed, 276
- macrophages, 277
- microphages, 277
- motile, 276
- Phagocytosis, 272
- acquired immunity and, 298
- alexin extraction in, from leukocytes and lymphatic organs, 304 *et seq.*
- chemotaxis in, 285
- influence of bacteria in, 288, 289
- influence of bacterial extracts in, 288
- malic acid in, 286
- of slime-molds or myxomycetes, 286
- of spermatozoa of ferns, 286
- Pfeffer's technique in, 286
- destruction of bacteria in, 297, 300
- by alexin (or cytase) in leukocytes, 301, 302
- action of, 302
- Metchnikoff's interpretation of, 302
- by exudates, 300
- by phagocytes, 300
- destruction of red blood cells by, 276
- differences in degree of, due to bacteria, 325
- differences in phagocytic energy in, due to leukocytes in, 324
- differences in virulence of bacteria, dependent on their resistance to leukocytes in, 325
- digestion among protozoa and, 274
- "dust cells" in, 279
- early investigations in, 272
- endothelial cells in, 278, 279
- enzymes in, endocellular and extracellular, 305
- eosinophile cells in, 278
- Phagocytosis, fixateur or sensitizer in, action of, in immunized animals, 301
- giant cells in, 280
- foreign body, 280
- tuberculous, 280
- in daphnia, 296
- in higher animals, 296
- in immune serum, 315
- bacteriolysins in, bactericidal sensitizers and, 321 *et seq.*
- heated, opsonic action in, increase of, by addition of fresh normal serum, 318
- increase of, 311
- attributed to "stimulins," 311
- with addition of leukocytes, 312
- opsonin contents a factor in, 313
- opsonins in, increase of, 315
- normal opsonins and, 320 *et seq.*
- specificity of, 321
- thermostability of, 320
- in immunity, 90
- in normal serum, opsonins in, coöperation of heat-sensitive body in, 319
- nature of, 316
- similarity of, to alexin, 316, 317
- specific thermostable, 317
- specificity of, 318
- in process of inflammation, 280 *et seq.*
- with pyogenic staphylococci, 281
- with tubercle bacilli, 283
- increase of, by injection of leukocyte extracts, 308 *et seq.*
- in increased resistance, 329
- with acquisition of immunity, 299
- intracellular digestion and, 274
- in vertebrates, 275
- leukocytes in, 324
- action of, 275, 276
- polynuclear, 278
- leukocytosis in, 290
- bacteria decreasing, 290
- bacteria increasing, 290
- lymphocytes in, large, 278
- measure of degree of, in active immunization, 329
- Leishmann's technique for, 329
- Simon, Lamar and Bispham's technique for, 332, 333
- value of, in therapeutics, 338
- Wright's technique for, 330 *et seq.*
- Phagocytosis, measure of degree of, Wright's technique for, difficulties in, 332, 333
- value of, 333
- mechanism of process of, 280 *et seq.*
- Metchnikoff's early investigations on, 273, 274
- normal and immune opsonic action in, quantitative differences between, 324
- normal degenerative and retrogressive processes and, 276
- observation of, *in vitro*, 313
- of micro-organisms, with or without culture media, 297
- opsonins in. *See* Opsonins
- phagocytes engaged in, varieties of, 276
- fixed, 276
- macrophages, 277
- microphages, 277
- motile, 276
- process of inflammation in, 280 *et seq.*
- proteolytic enzymes from leukocytes in, 306, 307
- qualitative difference between normal and immune opsonic substances in, 315, 316
- reduction of phagocytic activity in, 298
- by growth of bacteria within leukocytes, 298
- by protection of bacteria from phagocytes, 299
- by use of narcotics, 299
- relation of, to active immunity, 329
- relation of virulence to, 312
- removal of extravasations of blood and, 275
- resistance of bacteria to, due to non-absorption of opsonin, 326
- resistance of infected subject and, 296, 297
- resistance of virulent bacteria to, in normal serum, 325
- spontaneous, 313
- tissue cells in, 278
- varieties of body cells engaged in, 278
- dependent on nature of invading substance, 279
- Pick and Yamanouchi's experiments on two separate substances in anaphylactic antigen, 388
- work on anaphylaxis, 371

- von Pirquet and Schick's studies of serum sickness, 427 *et seq.*
- von Pirquet's tuberculin skin reaction, 440
- Placental cytotoxin, 92
- Plague, active prophylactic immunization against, 486
- Besredka's vaccines in, 487
- Haffkine's early work on, 486
- Kolle and Otto's investigations in, 487
- Rowland's vaccine in, 487
- Strong's investigations in, 487
- relative susceptibility of man and animals to, 53
- serum therapy of, 478
- Dean's serum in, 480
- Lustig's serum in, 480
- Markl's serum in, 479
- Rowland's serum in, 480
- value of, 480, 481
- Yersin, Calmette and Borrel's investigations in, 478
- Yersin's serum in, 478-480
- value of, 479
- "Plasmines," 72
- Pneumococcus infection, relative susceptibility of man and animals to, 54
- Pneumococci, mutation of, 472
- Pneumonia, agglutination reaction in diagnosis of, 221
- serum therapy in, Cole's work on, 475
- Dochez and Gillespie's work on, 475
- nature of action in, 474, 475
- Neufeld and Haendel's work on, 474
- Poison-ivy, specific antitoxin from, 96
- Pollantin, 435
- Poliomyelitis, relative susceptibility of man and animals to, 55
- Polyceptors (Ehrlich), 156
- Precipitation, 248
- inhibition zones in, 162
- Precipitin reaction, 248
- against heated proteins, 258 *et seq.*
- cotoprecipitin in, 258
- experiments on, 260-262
- heat-alkali-precipitin in, 259, 260
- native precipitin in, 259
- 70° precipitin in, 259
- Schmidt's experiments on, 260, 261
- agglutination reaction analogous to, 263
- analogy of, to colloidal flocculation, 265
- autocytotoxins in, 263
- Precipitin reaction, bacterial precipitins in, partial or minor, 252
- specificity of, 251, 252
- Ehrlich's conception of, 264
- electrolytes in, effect of, 265
- forensic blood test in, 257
- ring test of Fornet and Müller in, 257
- group reactions of bacterial precipitins in, 251, 252
- diagnostic value of, 252
- heat precipitins in, 259
- heated precipitating serum, effect of mixed sera in, 266-267
- protective action of, 266
- inhibition zones in, 265, 266
- isoprecipitins in, 255
- medico-legal value of, 254
- non-specific partial reactions in, elimination of, 254
- Nuttall's experiments on determining zoölogical classifications by, 254, 255
- organ specificity in, 262, 263
- precipitinogen in, 249
- chemical nature of, 249
- effect of heating on, 258
- non-protein, 249, 250
- obtaining of, 249
- precipitinoids in, formation of, 265
- precipitins in, delicacy of, 253
- determination of potency of, 253
- inactivation of, by heat, 264
- effect of, in bacterial filtrates, 264
- production of, against unformed proteins, 252, 253
- methods of, 251
- of specific, 249
- by pepton, 251
- effect of heating on, 249
- in animal sera by foreign protein, 248
- structure of (Ehrlich), 264
- zymophore group in, 264
- effect of heat on, 265
- quantitative proportions in, effect of, 265, 266
- relative concentration of reacting bodies a factor in, 265, 266
- residue antigen and antibody in, 267 *et seq.*
- explanations of, 268 *et seq.*
- Precipitin reaction, residue antigen and antibody in, experiment on, 269
- salts in, effect of, 265
- species determination by means of, 253, 254
- species specificity in, 262
- specificity of, 248, 253, 254
- vegetable proteins determined by, 255
- zoölogical classifications by means of, 254, 255
- Precipitin tests, methods of performing, 255 *et seq.*
- forensic blood test in, 257
- ring test of Fornet and Müller in, 257
- Precipitinogen, 249
- chemical nature of, 249
- effect of heating on, 258
- non-protein, 249, 250
- nature of, 250
- obtaining of, 249
- Precipitinoids, 265
- Precipitins, 248
- against heated proteins, 258 *et seq.*
- cotoprecipitin, 258
- experiments on, 260-262
- heat-alkali-precipitin, 259, 260
- native precipitin, 259
- 70° precipitin, 259
- Schmidt's experiments on, 260, 261
- bacterial, group reactions in, 251
- partial or minor, 252
- specificity of, 251, 252
- definition of, 90
- delicacy of, 253
- determination of potency of, 253
- heat, 259
- in hemolytic serum, 94
- inactivation of, by heat, 264
- effect of, in bacterial filtrates, 264
- isoprecipitins, 255
- organ specificity of, 262, 263
- production of, 129, 249
- against unformed proteins, 252, 253
- methods of, 251
- specific, by pepton, 251
- effect of heating on, 249
- in animal sera by foreign protein, 248
- "species," specificity of, 262, 263
- specific, 248
- discovery of, by Rudolf Kraus, 248
- structure of (Ehrlich), 264
- zymophore group in, 264
- effect of heat on, 265
- Pregnancy, diagnostic value of Abderhalden's protective ferments in, 496

- Pro-agglutinoïd phenomenon in agglutination explained as protective colloid action, 236
- Pro-agglutinoïd zone in agglutination, 162
- complement deviation reaction, analogous to, 162
- Pro-agglutinoids, 235
- Prophylactic immunization, a active, in man, 481
- against cholera, 484.
See also under Cholera
- against plague, 486. *See also under* Plague
- against rabies, 489. *See also under* Rabies
- against small-pox, 488. *See also under* Small-pox
- against typhoid fever, 482. *See also under* Typhoid fever
- "Protection," 195
- "Protein fever," 367
- Proteins, unknown, alexin fixation test in determination of nature of, 211
- delicacy of, 213
- technique of, 212
- Prototoxoids, 115
- Protozoa, digestion among, and its relation to phagocytosis, 274
- Ptomains, as indirect cause of infection, 30
- chemistry of, 29
- definition of, 31
- relation of, to infection, 28
- Putrefaction, chemistry of, 29
- micro-organisms causing, 1
- Pyemia, 25
- Rabies, active prophylactic immunization against, 489
- Högyes method of treatment in, 492
- Pasteur's work on, 489
- preparation and attenuation of virus for, 490
- treatment of patients in, 491
- Ranzi's work on anaphylaxis, 367
- Rattlesnake poison, antitoxin for, 466
- Receptors, cell, overproduction of, 152
- complementophile, 149
- cytophile, 149
- definition of, 126
- of third order, 150
- sessile, in anaphylaxis, 290
- Resistance. *See also* Immunity
- acquired, 60
- bactericidal properties in serum and, 297
- body temperature and, 51
- cellular theory of, 80
- Resistance, cultural conditions for bacteria in body and, 56
- degree of phagocytosis and, 296, 297
- humoral theory of, 80
- increased invasive powers of bacteria and, 56
- individual differences and, 58
- inheritance a factor in, 56
- local, in skin infections, 102
- natural, against infection, 49
- racial differences in, 55
- species, to infection, 51
- "Retention theory of immunity," 83
- Rhus toxicodendron, specific antitoxin from, 96
- Richet and Héricourt's work on anaphylaxis, 360
- Richet and Portier's work on anaphylaxis, 360
- Richet's work on passive anaphylaxis, 380
- Ricin, "protein-free," 96
- Römer's method for diphtheria antitoxin standardization, 461
- "Root-tubercle" bacilli, 7
- Rosenau and Anderson, researches of, in bacterial anaphylaxis, 410
- work on anaphylaxis, 362, 374 *et seq.*
- Rosenow on variations in streptococci, 472
- Roux and Yersin, experimental immunization in hog cholera by, 72
- Rowland's antiplague serum, 480
- vaccine in prophylactic immunization against plague, 487
- Russell's vaccines for prophylactic immunization against typhoid fever, 483
- Salt-inactivation of alexin, 178
- Salts, effect of, in agglutination, 243
- in precipitation, 265
- Saprophytes, biological transition of, to parasites, 5
- occurrence of, 2
- pathogenic powers of certain, 4
- pure, 11
- Saprophytic micro-organisms, definition of, 4
- Scarlet fever, relative susceptibility of man and animals to, 54
- Schmidt, precipitation experiments of, on heat precipitins, 260, 261
- Sensitization, 359
- Bordet's views on, 162, 163
- complement deviation in, 160
- Ehrlich and Sachs' phenomenon in, 165
- Bordet and Gay's interpretation of, 166, 167
- Ehrlich and Sachs' views on, 164, 165
- Neisser-Wechsberg phenomenon in, 160
- Septicemia, chronic, adaptation of bacteria in, 7
- secondary foci in, 7
- Sensibilisin, 387
- Sensibilisinogen, 387
- Sensitized bacteria, immunization with, 68
- Sensitized tuberculin, 357
- Sensitized vaccines, 355
- Sensitizer. *See* Amboceptor
- Bordet's definition of, 159
- quantitative determination of, in immune serum, 160, 161
- Serum. *See also* Blood serum
- antitoxic, direct effect of, on toxin, 104
- indirect protective action of, against toxin, 104
- bactericidal properties of, in immunity, 81
- resistance and, 297
- cell-free, bacterial growth in, 81
- immune, bacteriotropins in, without lysins, 322
- heated, reactivation of, by addition of alexin, 318
- opsonins in, bactericidal sensitizers and, 321 *et seq.*
- heated, increase of power of, by addition of fresh normal serum, 318
- increase of, 315
- normal opsonins and, 320 *et seq.*
- specificity of, 321
- phagocytosis in, 91, 315
- increase of, 311. *See also* Phagocytosis
- normal, agglutinins in, 91
- an anticomplementary properties of, 196
- hemolytic properties of, 91
- opsonic powers of, 314
- reduction of, by heat, 314
- opsonins in, 91
- coöperation of heat-stable and heat-sensitive body in, 318, 319
- nature of, 316

- Serum, normal, opsonins**
in, resistance of,
to heat, 315
similarity of, to
alexin, 316, 317
specificity of, 318
thermostability of,
320
specific thermostable
opsonins in, 317
normal antitoxic, diph-
theria, 107
opsonins in normal and
immune, qualita-
tive differences in,
315, 316
- Serum sickness, 426**
analogy of anaphylaxis
with, 428
antibody formation in,
429
incubation time in,
429
methods of administra-
tion of antitoxin
to avoid, 430
Besredka's method of,
431
by alteration of serum,
430
Friedberger and Mita's
method of, 432
in animal experimen-
tation, 431
with concentrated an-
titoxin, 430
von Pirquet and Schick's
studies of, 427 *et*
seq.
symptoms of, 426
accelerated reaction of
von Pirquet and
Schick in, 427
after first injection,
426
after second injection,
427
immediate reaction in,
427
analogy of, to ana-
phylaxis, 427
- Serum therapy, anaphylax-
is in. See Serum
sickness**
in diphtheria, 446. *See*
also Diphtheria
antitoxin
in diseases caused by
bacteria which do
not form soluble
toxins, 466
action of serum upon
extensive infec-
tion in, 467
antibacterial action in,
466
in epidemic cerebrospinal
meningitis, 469.
See also under
Cerebrospinal
meningitis, epi-
demic
in plague, 478. *See also*
Plague, serum
therapy of
in pneumonia, 474. *See*
also Pneumonia,
serum therapy
in
in streptococcus infec-
tions, 471. *See*
also under Streptococcus
infections
- Serum therapy, in typhoid
fever, 475. See**
also Typhoid
fever, serum ther-
apy of
- Side chain theory, anti-
body production
in body cells in,
130**
body cell in, 125
chemical nature of,
126
"Leistungskern" in,
126
side chains or recep-
tors in, 126
chemical action of anti-
gens in, 128
definition of side chains
in, 126
diagram showing cell-
receptors and im-
mune bodies (Ehr-
lich) in, 127
in toxin-antitoxin reac-
tion, 124
overproduction of recep-
tors in, 129
flow of, into blood a
cause of immu-
nity, 128
physical mechanism of,
124
recapitulation of, 130
relationship between
susceptibility of
tissue and toxin-
binding properties
in, 131
- Skin, defence of, in infec-
tion, 12**
- Skin infections, local im-
munity in, 102**
- Small-pox, active prophylac-
tic immuniza-
tion against,
488**
Jenner's discovery of,
488
production and prepa-
ration of vac-
cine for, 488
history of experimenta-
tion in immuniza-
tion against, by
Jenner, 62
relative susceptibility of
man and animals
to, 54
vaccination for, history
of, 62
principles of, 62
- Smith, Theobald, investiga-
tions of, in ana-
phylaxis, 363**
phenomenon of, in ana-
phylaxis, 361
- Snake venoms, 36**
action of, 465
activation of, by endo-
complement in
blood cells, 174
by sera, 174
antitoxins for, 464
effect of heat on, 105
immunization against,
464
Kyes' experiments in,
174, 175
neurotoxins in, 465
peculiarities of, 464
toxin-antitoxin combina-
tion in, stability
of, 105
- Snake venoms, toxin-HCl
modification of,
effect of heat on,
106**
toxin-antitoxin reaction
with, 105, 465
filtration experiments
in, 105
neutralization theory
of, 105
time element in,
105
- Sols, 500**
Species resistance, 51
Specificity, definition of, 76
Spermatotoxins, 92
Spleen, removal of, and
antibody forma-
tion in active im-
munity, 100
and susceptibility to
infection, 101
- Standardization of anti-
toxin, guinea pigs
used in, 108**
minimum lethal dose in,
109
- Standardization of diph-
theria antitoxin,
by means of tox-
in, 107**
early attempts at, 107
unit in, 107
- Standardization of sera,
Pfeiffer's method
of, 139**
- Standardization of tetanus
antitoxin by
means of toxin,
107**
- Standardization of vac-
cines, 352**
Hopkins' method of,
353
Wright's method of, 352,
353
- Staphylococcus furunculo-
sis, opsonic index
in vaccine treat-
ment of, 335**
- Staphylococcus infections,
opsonic index in,
334**
during vaccine treat-
ment with dead
staphylococcus
cultures, 335
relative susceptibility of
man and animals
to, 54
- Stern's modification of
Wassermann test,
209**
- Stimulins, 311**
- Strauss test, 25**
- Streptococci, variations in,
472**
- Streptococcus infections,
agglutination re-
action in diag-
nosis of, 222**
relative, of man and
animals, 54
serum therapy in, 471
difficulties in, owing
to variations of
streptococci, 471,
472
early investigations in,
472
Marmorek's work on,
472, 473
nature of action in,
473

- Streptococcus infections, serum therapy in, standardization of serum in, 474
value of, 473
- Strong's investigations in prophylactic immunization against plague, 437
method of prophylactic vaccination in cholera, 436
- Sub-infection, 24
- "Summation of negative phase" in active immunization, 338, 339
- "Summation of positive phase" in active immunization, 339
- Surface tension in chemotaxis, 293
- Susceptibility, body temperature and, 51
cultural conditions for bacteria in body and, 56
increased invasive powers of bacteria and, 56
individual differences and, 58
inheritance a factor in, 56
racial differences in, 55
relative, of animals to infection, 51
species resistance to infection and, 51
- Sycosis, opsonic index in vaccine treatment of, 336, 337
- Syntoxoids, 116
- Syphilis, diagnosis of, B o r d e t - G e n g o u phenomenon in, 188
by alexin fixation, 198, 199
by direct precipitation of syphilitic serum by emulsions of lecithin and of sodium glycocholate, 204
relative susceptibility of man and animals to, 52
ultramicroscopic method of finding precipitates in sera of, 204
- Wassermann reaction in, diagnostic value of, 210, 211
technique of, 207
Bauer's modification of, 209
Noguchi's modification of, 208, 209
schematic presentation of, 209
Stern's modification of, 209
- Temperature, body, and resistance to infection, 51
- Tetanus, "cryptogenic," 5
relative susceptibility of man and animals to, 53
- Tetanus, toxin fixation in, by brain tissues, 131
lipoidal, substances a factor in, 133
proteolytic enzymes a factor in, 133
temperature a factor in, 132
- Tetanus antitoxin, production of, 463
standardization of, 463
by means of toxin, 107
- Tetanus bacillus, action of, 4, 5
- Tetanus toxin, action of, 41
- Thread reaction of Pfaundler in agglutination, 222, 223
- Thymotoxin, 92
- Thyroid gland, production of alexin in, 172
production of opsonins in, 173
- Toxemia, 10
- Toxicity, definition of, 11
- Toxin-antitoxin, diphtheria. *See* Diphtheria toxin-antitoxin
- Toxin-antitoxin combination, chemical relations of, 114
effect of heat on, 106
in snake venom, stability of, 105
toxin-HCl modification of, 106
effect of heat on, 106
measurement of, by partial absorption method of Ehrlich, 115
stability of, 105
valency of component parts of, 114
- Toxin-antitoxin reaction, analogy between chemical reactions and, 118, 119
antibody production in body cells in, 130
body cell in, 125
chemical nature of, 126
chemical action of antigens in, 128
concentration of reagents in, 107
degrees of toxicity in, 123
direct neutralization the protective power of, 124
effect of heat on, 105
effect of temperature on, 104
mechanism of, 104
neutralization in, absorption theory of, 123
Arrhenius and Madsen on, 126
Bordet on, 122
Bordet - Danysz phenomenon in, 123
von Dungern's views on, 124
Danysz effect in, 123
phenomena of, 119 *et seq.*
- Toxin-antitoxin reaction, overproduction of receptors in, 128
flow of, into blood a cause of immunity, 128
physical mechanism of, 124
quantitative relations in, 106, 123
relationship between susceptibility of tissue and toxin-binding properties in, 131
side chain theory in, 124
specificity of, 124, 129
speed of action of, 107
time element in, 105
with snake venom, 105
filtration experiments in, 105
neutralization theory of, 105
time element in, 105
- Toxin, bacterial. *See* Bacterial toxins.
chemical relations of, with antitoxin, 114
deterioration theory of, 110
definition of, 32
differences in combining avidity of, 111
diphtheria, construction of, 118
normal, 107
direct effect of antitoxic serum on, 104
epitoxoid form of, 112
indirect effect of antitoxic serum on, 104
structure of, 110
toxoid, 110
prototoxoids in, 115
syntoxoids in, 116
toxin, 113
true, 33
analogy of, with enzymes, 36
bacteria producing, 34
characteristics of, 34
chemically indefinable nature of, 35
diseases for which some investigators claim, 469
heat sensitiveness of, 36
incubation time of, 36
production of antitoxin by, 35
- Toxin hypersusceptibility, anaphylaxis and, 407
- Toxin spectra, construction of, 116
definition of, 116
measurement of, 116, 117
principles of, 116
- Toxin unit, definition of, 109
- diphtheria, 107
- Toxoids, definition of, 110
- Toxons, action of, 113
definition of, 113
structure of, 113
- Toxophore group of toxin, 110
action of, 110

- Tubercle bacilli, effect of body temperature on virulence of, 12
- Tuberculosis, avian type, relative susceptibility of animals to, 52
- bovine type, relative susceptibility of man and animals to, 52
- human type, relative susceptibility of man and animals to, 52
- meistagmin reaction in diagnosis of, 497
- of cold-blooded animals, immunity of warm-blooded animals to, 52
- opsonic index in, 341-342, 343
- Tuberculin ophthalmoreaction, anaphylactic nature of, 440
- Tuberculin reaction, analogy of, to anaphylaxis, 442
- anaphylactic nature of, 438
- Ball's experiments with passive sensitization in, 443
- diagnostic value of, 442
- Koch's experiments in, 439
- nature of, 438
- Babes' interpretation of, 439
- Koch's interpretation of, 439
- Wassermann and Brucks' interpretation of, 439
- specific antibody formation in, 442
- Tuberculin skin reaction, anaphylactic nature of, 440
- von Pirquet's interpretation of, 441
- Tuberculins, 355
- Bouillon Filtre (Denys), 357
- New Tuberculin (TR and TO), 356
- New Tuberculin Bacillary Emulsion, 357
- Old Tuberculin (Koch), 355
- Sensitized Tuberculin, 357
- Tumors, malignant, alexin fixation in diagnosis of, 213
- von Dungern's method of, 214
- antigen production for, 214
- results of, 215
- technique of, 214 *et seq.*
- organ-specific qualities in, 373
- Typhoid bacilli, attenuation of virulence of, 18
- Typhoid carriers, 3
- Typhoid fever, adaptation of bacteria in, 8
- agglutination reaction for diagnosis of, 219
- Typhoid fever, agglutination reaction for diagnosis of, microscopic method, 219
- microscopic method, 220
- effect of path of introduction of bacteria of, on infection, 14
- meistagmin reaction in diagnosis of, 497
- prophylactic immunization against, active, 482
- early experimentation in, 482
- living sensitized vaccines used in, 484
- results of, in United States Army, 483
- Russell's vaccines in, 483
- sensitized killed vaccines in, 484
- relative susceptibility of man and animals to, 54
- serum therapy in, 475
- Besredka's anti-endotoxic serum in, 476
- Chantemesse's early experiments in, 475
- Garbat and Meyer's work on, 477
- Kraus and Stenitzer's serum in, 477
- Gottstein - Mathes' work on, 477
- Lüdke's work on, 477
- nature of reaction in, 476
- Typhus fever, relative susceptibility of man and animals to, 54
- Ultramicroscope, 503
- Vaccination, prophylactic, in man, 481
- in anthrax, 64
- in cholera, 484. *See also under Cholera*
- in plague, 486. *See also under Plague*
- in rabies, 489. *See also under Rabies*
- in small-pox, 488. *See also under Small-pox*
- history and general principles of, 62
- in typhoid fever, 482. *See also under Typhoid fever*
- Vaccine therapy. *See also Immunization, active*
- anaphylaxis in, 432
- as a therapeutic measure, action of, in local infections, 346
- in generalized systemic infections, 347
- in successive local infections, 347
- value of, in acute diseases, 350
- Vaccine therapy, as a therapeutic measure, value of, in subacute or chronic cases, 349
- autoinoculations by massage or exercise in, 340
- "high tide" of immunity in, 340
- "negative" phase in, 338
- second injection in, 338
- successive inoculations in, 338, 339
- summation of, 338, 339
- opsonic index in, 328 *et seq.*
- comparison between that in exudate of infected foci and blood serum, 340
- improvement and, 341
- in tuberculosis, 341-343
- Leishmann's technique for determination of, 329
- Simon, Lamar and Bispham's technique for determination of, 332, 333
- value of, 338
- in controlling therapeutic vaccinations, 344
- in showing degree and conditions in which vaccination is successful, 344, 345
- Wright's technique for determination of, 330 *et seq.*
- difficulties in, 332, 333
- value of, 333
- relation of phagocytosis to, 329
- second positive phase in, 339
- "summation of positive phase" in, 339
- tuberculins in, 355
- value of, as prophylactic measure, 345, 346
- as therapeutic measure, 346
- Vaccines, autogenous, 351
- production of, 351
- with dead bacteria, 351
- with living bacteria, 351
- sensitized, 355
- standardization of, 352
- Hopkins' method of, 353
- Wright's method of, 352, 353
- Vaughan's work on anaphylaxis, 366, 367
- on bacterial anaphylaxis, 412
- Vaughan and Wheeler, proteid split products of, in anaphylactic poison, 403
- theory of, on mechanism of anaphylaxis, 393

- Vaughan and Wheeler, work of, on toxic fraction of protein molecule in anaphylaxis, 393
- Virulence, aggrassin secretion of bacteria in body and, 20-22
- capsule formation of bacteria and, 18
- definition of, 11
- dependent on resistance of bacteria to leukocytes in phagocytosis, 325
- ectoplasmic hypertrophy of bacteria in relation to, 19, 20
- effect of body temperature on, 12
- effect of cultural adaptation of bacteria on, 12
- effect of path of introduction of bacteria on, 12-14
- effect of quantity of bacteria introduced on, 14
- increase of, by attenuation of bacteria, 17
- measurement of relative degrees of, 15
- of capsulated bacteria, 326
- relation of, to phagocytosis, 312
- relative to number of bacteria introduced, 15
- specificity of bacteria and, 22
- variation in, of bacteria successively passed through animals, 16, 17
- of different strains of same bacteria, 15, 16
- Virulins, 22, 326
- "Virus fixe" in treatment rabies, 490
- Wassermann reaction, 198
- alexin fixation principle in, 198
- not by union of specific syphilitic antigen with spirochæta pallida antibodies, 204
- alexin titration in, 206
- antigen preparation for, 200
- by addition of cholestrin, 201
- by method of Brownling and Cruikshank, 201
- by method of Noguchi, 200
- by methods of Porges and Meier, 200
- by methods of Weil and Braun, 200
- titration in, 202
- diagnostic value of, 210, 211
- in diagnosis of syphilis, 198
- in diseases other than syphilis, 210
- in normal organs, 200
- Klausner theory in, 204
- precipitation in, by addition of syphilitic serum to lecithin emulsions, 204
- produced with syphilitic serum in antigens from normal organs, 200
- specific antigen from spirochæta pallida cultures unsuitable in, 203
- Wassermann reaction, spinal fluid used in performance of, 210
- technique of performance of, 207
- Bauer's modification of, 209
- Noguchi's modification of, 208, 209
- schematic presentation of, 209
- refrigerator method in, 208
- Stern's modification of, 209
- schematic presentation of, 207
- theories of, 204, 205
- titration of hemolytic amboceptor or sensitizer in, 205
- ultramicroscopic method of finding precipitates in syphilitic sera in, 204
- Weigert's law of overcompensation, 128
- Wright's method of standardization of vaccines, 352, 353
- Wright's technique for determination of opsonic index, 330 *et seq.*
- difficulties in, 332, 333
- value of, 333
- Wright's studies of bactericidal and agglutinating powers of blood serum, 328
- Yellow fever, susceptibility to, 55
- Yersin anti-plague serum, 478-480

THE following pages contain advertisements of a few of the
Macmillan books on kindred subjects.

THE CANCER PROBLEM

By WM. SEAMAN BAINBRIDGE, M.D.

Surgeon to the New York Skin and Cancer Hospital

Illustrated with many rare plates. Price, \$3.50 net.

Dr. Bainbridge's experience in the diagnosis and treatment of malignant diseases, and his familiarity with the experimental investigation of the disease, have enabled him to exercise rare discrimination in the selection of the subject matter for his forthcoming book. Great care has also been exercised in its presentation and in the preparation of its beautiful microscopic plates, of which there are a large number.

The Cancer Problem will prove useful alike to the laboratory worker, to the general practitioner, to the surgeon, to the student of public hygiene and health—in fact, to all who are interested in matters of general and individual health-maintenance. It is a practical summary of what is known today concerning cancer.

TABLE OF CONTENTS

DEDICATION.

PREFACE.

SECTION I. HISTORY.

SECTION II. GENERAL DISTRIBUTION: CHAPTER 1—BOTANICAL DISTRIBUTION.

CHAPTER 2—ZOOLOGICAL DISTRIBUTION. CHAPTER 3—GEOGRAPHICAL AND ETHNOLOGICAL DISTRIBUTION.

SECTION III. STATISTICAL CONSIDERATIONS.

SECTION IV. ETIOLOGY: 1—THEORIES: (A) Early Theories; (B) Theories which have engaged attention since the beginning of Modern Cancer Research. 2—PRE-DISPOSING CAUSES.

SECTION V. HISTO-PATHOLOGY.

SECTION VI. CANCER RESEARCH—A RESUME OF THE WORLD'S WORK.

SECTION VII. CLINICAL COURSE AND DIAGNOSIS: 1—CLINICAL COURSE. 2—DIAGNOSIS. 3—POSSIBLE ERRORS IN DIAGNOSIS: (A) Head; (B) Tongue; (C) Breast; (D) Abdomen; (E) Pelvis.

SECTION VIII. PROPHYLAXIS.

SECTION IX. THE INVESTIGATION OF "CANCER CURES."

SECTION X. NON-SURGICAL TREATMENT: 1—CAUSTICS OR ESCHAROTICS.

2—PHYSIO-THERAPY: (A) Heat; (B) Light; (C) Electricity; (D) Radioactivity.

3—BIO-THERAPY: (A) Serotherapy; (B) Vaccinotherapy; (C) Opothrapy.

SECTION XI. SURGICAL TREATMENT: 1—GENERAL TECHNIC OF SURGERY AS

APPLIED TO CANCER. 2—SPECIAL TECHNIC: A. SURGICAL TECHNIC AS MODIFIED IN ACCORDANCE WITH THE "PERMEATION THEORY" OF THE DISSEMINATION OF CANCER.

B. PLASTIC PROCEDURES. C. PALLIATIVE PROCEDURES: (1) *Nervous System*—(a)

Neurectomy; (b) Decompression. (2) *Lymphatic System*—(a) Lymphangioplasty;

(b) Paracentesis Abdominalis; (c) Thoracocentesis. (3) *Vascular System*—Arterial

Ligation. (4) *Respiratory System*—Tracheotomy. (5) *Alimentary System*—(a)

Esophagostomy; (b) Gastrostomy; (c) Gastroenterostomy; (d) Colostomy; (e)

"Short-Circuit"; (f) Colectomy. (6) *Urinary System*—(a) Cystostomy; (b) Neph-

rotomy; (c) Ureteral Transplantation. (7) *Biliary System*—(a) Cholecystostomy;

(b) Cholecystenterostomy.

SECTION XII. IRREMOVABLE CANCER.

SECTION XIII. INSTITUTIONS FOR THE CARE OF CANCER PATIENTS.

SECTION XIV. CAMPAIGN OF EDUCATION.

THE OUTLOOK.

GENERAL BIBLIOGRAPHY.

INDEX OF AUTHORS. INDEX OF SUBJECTS.

PUBLISHED BY

THE MACMILLAN COMPANY

64-66 Fifth Avenue, New York

Researches on Rheumatism

By

F. J. POYNTON, M. D.

London, Vice-Dean of University College Hospital Medical School, Senior Physician to
Cut-patients at University College Hospital, London, etc., and

ALEXANDER PAINE, M. D.

London, D.P.H. England, Director of the Cancer, Research Institute, London.

With frontispiece in color and 106 illustrations, 8vo, \$5.00 net.

This collection of papers is the result of fifteen years' work by these two noted authorities, who are well known to the American profession for their researches in this subject. This product of their joint authorship makes a remarkable history of achievement and steady advance in our knowledge, and it is needless to say that the literary quality is of the highest. The book unquestionably gives the most recent and authentic presentation of a very important group of diseases and at the conclusion of the volume the bearing of these investigations upon clinical medicine and public health is considered in a special article. Etiology, pathology, symptomology, diagnosis, prognosis, treatment and prevention are summarized at the end, so that this work stands as a complete treatise on the subject. The illustrations have been chosen with the intention of demonstrating the intimate processes of rheumatism in the body, and to act as a pictorial guide to the main conclusions. Their workmanship is so perfect that they are almost equal to the microscopic slide itself.

"The papers are for the most part arranged in chronological order, and taken together form a consistent account of a very protracted research which has acquired a special value as the joint work of clinicians and bacteriologists. . . . The record of work that they have now put forth will doubtless remain as a valuable landmark in the course of discovery."

—*British Medical Journal.*

PUBLISHED BY

THE MACMILLAN COMPANY

64-66 Fifth Avenue, New York

Manual of Bacteriology

By

ROBERT MUIR, M.A., M.D., F.R.C.P., Ed.

Professor of Pathology, University of Glasgow

AND

JAMES RITCHIE, M.A., M.D., B.Sc.

Reader in Pathology, University of Oxford

New American Edition. Revised and Enlarged. \$3.25 net.

"The American Edition of this well-known Manual is perhaps one of the best and most comprehensive, up-to-date hand-books for the student published in the English Language. The treatment of the doubtful questions is to be commended. The investigations of each observer and the conclusions are stated with as little bias as possible.

. . . . An appendix furnishes a compact outline of the principal literature on the different subjects. This outline deals chiefly with the original works found in the foreign languages."—*Medical Record*.

"Like Gray's Anatomy, Green's Pathology, Parke's Hygiene, and other classical text-books, this manual is destined to remain for years to come the favorite of both teacher and student, to whose needs it is so admirably adapted."—*Philadelphia Medical Journal*.

"A very useful work for the purpose intended."

—*International Medical Magazine*.

PUBLISHED BY
THE MACMILLAN COMPANY
64-66 Fifth Avenue, New York

Tuberculosis of Bones and Joints in Children

By John Fraser, M. D., F. R. C. S. (Edin.), Ch. M., Assistant-Surgeon, Royal Hospital for Sick Children, Edinburgh. Containing 34 full page illustrations—4 of them in color—and many line drawings in the text.
Crown, 8vo., cloth, \$4.50 net.

Tuberculous disease of the bones and joints is in large measure a disease of children. This work is intended as an exposition of this subject. Our recent knowledge of the cause of the disease has been discussed, and the author's well-known investigations into the pathology have been incorporated in the book. The treatment of the various conditions is thoroughly dealt with, and the illustrations are a special feature. The work is essentially intended for the practical information of students and practitioners.

A Manual of Immunity

By Elizabeth T. Fraser, M. D. (Glas.) Late Assistant Bacteriologist, Glasgow Royal Infirmary.
Cloth, 192 pp., index, \$1.60 net.

This book is intended primarily for the busy general practitioner whose lack of time forbids laboratory work. During the last twenty years, the science of Immunity has been the most living branch of Medicine, and in these days of Sero-diagnosis and Vaccine and Serum Therapy, it is obvious that the physician can no longer be a mildly interested on-looker. This Manual, therefore, supplies in a thorough manner the fundamental facts of Immunity.

On Diseases of the Rectum and Anus

By Harrison Cripps, F. R. C. S., Consulting Surgeon, St. Bartholomew's Hospital. Fourth Edition, including the Sixth Edition of the Jacksonian Prize Essay on Cancer, and the Opening Address on the Surgical Treatment on Rectal Cancer, delivered at the Annual Meeting of the British Medical Association, Liverpool, 1912.

Illustrated. Cloth, 8vo, \$3.25 net.

Dr. Cripps, the author of this important monograph, is well known in this country as an authority on these subjects and as a wide contributor to many American medical magazines. The clinical cases recorded in this monograph are largely drawn from his notes made in the Registers of St. Bartholomew's Hospital, and the pathological observations have been verified by postmortem or microscopic investigation.

PUBLISHED BY
THE MACMILLAN COMPANY

64-66 Fifth Avenue, New York

The Common Bacterial Infections of the Digestive Tract and the Intoxications Arising from Them.

By the late C. A. Herter, M.D., Professor of Pharmacology and Therapeutics in Columbia University; Consulting Physician to the City Hospital, New York.

Cloth, 8vo, 360 pages, \$1.50 net.

"The methods of investigating the digestive tract outlined in this volume will prove valuable to the practitioner who would determine the presence of abnormal bacterial processes before the onset of the clinical signs of incurable or highly refractory states of intoxication—and this is the all-sufficient *raison d'être* for the book. Considerable stress has been laid by the author on methods developed in his laboratory, in the belief that their painstaking application will furnish practitioners with reliable indications as to the progress of many cases of infection of the digestive tract."—*Merck's Archives*.

Practical Pathology, Including Morbid Anatomy and Post-Mortem Technique.

A Text-Book for Students and Practitioners. By James Miller, M.D., F. R. C. P. E., Assistant Pathologist to the Edinburgh Royal Infirmary; Lecturer on Pathology in the School of Medicine of the Royal Colleges, Edinburgh.

Small crown 8vo, cloth, ill., \$2.50 net.

This work is intended to give the student or practitioner in a compact form all necessary information relating to pathology in its more practical aspects. In dealing with the subjects of post-mortem technique, methods of fixing, staining, mounting of specimens, etc., an endeavor is made to give the student one reliable method in every case rather than puzzle him with numerous alternative methods. At the same time the different methods of fixing, staining, etc., adopted for special purposes, are not omitted. The book is illustrated with numerous drawings from photographs and actual specimens.

PUBLISHED BY
THE MACMILLAN COMPANY
64-66 Fifth Avenue, New York



